

BIONETICS

Summary of Mutagenicity Screening Studies Host-Medicated Assay Cytogebetics

Dominant Lethal Assay Contracy FDA 71-268 Compound FDA 71-36 Calcium

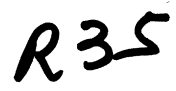
Propionate

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SUMMARY OF MUTAGENICITY
SCREENING STUDIES
HOST-MEDIATED ASSAY
CYTOGENETICS
DOMINANT LETHAL ASSAY
CONTRACT FDA 71-268
COMPOUND FDA 71-36
CALCIUM PROPIONATE

5516 Nicholson Lane Kensington, Maryland 20795

LBI PROJECT #2446



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SCREENING STUDIES
HOST-MEDIATED ASSAY
CYTOGENETICS
DOMINANT LETHAL ASSAY
CONTRACT FDA 71-268
COMPOUND FDA 71-36
CALCIUM PROPIONATE

SUBMITTED TO

FOOD & DRUG ADMINISTRATION
DEPARTMENT OF HEALTH, EDUCATION AND WELFARE
ROCKVILLE, MARYLAND

SUBMITTED BY

LITTON BIONETICS, INC. 5516 NICHOLSON LANE KENSINGTON, MARYLAND

MARCH 7, 1973

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March 7, 1973 September 13, 1974 - Revised

Mr. Leonard Appleby, Contracting Officer Department of Health, Education and Welfare Public Health Service Food and Drug Administration, CA-212 5600 Fishers Lane, Room 5C-13 Rockville, Maryland 20852

Reference: Contract FDA-268; LBI Project #2446

Dear Mr. Appleby:

Litton Bionetics, Inc., is pleased to submit a report for the referenced contract entitled "Mutagenicity Screening Studies" for compound FDA 71-36, Calcium Propionate.

Included in this report are the results and raw data of the three tests conducted: Host-Mediated Assay; Cytogenetic Studies; and Dominant Lethal Assay. Eight (8) copies are being submitted for your review.

Upon completion of the toxicology work an evaluation was made of our results to those appearing in the literature. In cases were our values were lower, the toxicology was repeated. In some instances either the Host-Mediated Assay, Dominant Lethal Assay, and/or Cytogenetic Studies were also repeated at one or more levels to fulfill the requirements of the contract. In some cases, the acute and/or subacute assays were involved.

If there are any questions concerning this report, or, if additional information is required, please do not hesitate to contact us.

Sincerely yours,

LITTON BIONETICS, INC.

David P. A. Fabrizig

Principal Investigator

DPAF:11s Enclosures (8)

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I. REPORT

A. <u>Introduction</u>

Litton Bionetics, Inc. (LBI) has investigated the possible mutagenicity of compounds selected and provided by the Food and Drug Administration under Contract 71-268. LBI's investigation utilized the three mammalian test systems herein described -- Host-Mediated Assay, Cytogenetic Studies and Dominant Lethal Assay. These tests provide information as to the types of genetic damage caused by environmental compounds -- pesticides, chemicals, food additives, drugs and cosmetics.

The Host-Mediated Assay is based upon the assumption that the action of a mutagen on the genetics of bacteria is similar to that in man.

This is further strengthened by the use of an eukaryotic organism (Saccharomyces cerevisiae). Since the mutation frequencies are well established for the indicator organism, any deviation due to the action of the test compound is readily detectable. As some compounds are mutagenic in bacteria and not in the host animal, and vice versa, this test is able to differentiate an action which may have been due to hosts' ability to detoxify or potentiate a suspected mutagen. This action is dependent upon the ability of the compound to gain access to the peritoneal cavity. Coupled with the direct action of the compound on the indicator organism in vitro, the assay provides a clear insight into host-mediation of mutagenicity.

Cytogenetics provides a valuable tool for the direct observation of chromosomal damage in somatic cells. Alteration of the chromosome number and/or form in somatic cells may be an index of mutation. These studies utilized examination of bone marrow cells arrested in C-metaphase from rats exposed to the test compound as compared to positive and negative control animals. If mutational



changes occur, the types of damage expected due to the action of chemicals are structural rearrangements, breaks and other forms of damage to the chromosomal complement of the cells exposed.

For the <u>in vitro</u> cytogenetic studies, we have a more rapid and inexpensive means of determining chromosomal damage. This is accomplished by observing cells in anaphase. As the chromatids separate and move along the spindle, aberrations may occur. Chromatids which do not migrate to the daughter cells may lead to uneven distribution of parts or of entire chromatids (mitotic nondysjunction). These give rise to "side arm" bridges which have been interpreted as point stickiness or localized failures of chromosome duplication point errors. These aberrations (bridges, pseudochiasmata, multipolar cells, acentric fragments, etc.) are extremely sensitive indicators of genetic damage.

The Dominant Lethal Test is an accurate and sensitive measure of the amount and type of fetal wastage which may occur following administration of a potential mutagen. Dominant lethal mutations are indicators of lethal genetic lesions. The effects of mutagens on the chromosomal complement of the spermatozoa of treated males results in alterations of form and number of chromosomes. Structural rearrangements and aneuploidy may lead to the production of non-viable zygotes, early and late fetal deaths, abortions and congenital malformations. In addition, aberrations could lead to sterility or reduced reproductive capacity of the F_1 generation. The action of a mutagen on specific portions of spermatogenesis is also apparent in this test.

B. <u>Objective</u>

The purpose of these studies is to determine any mutagenic effect of the test compound by employing the Host-Mediated Assay, Cytogenetic Studies



and the Dominant Lethal Assay, both <u>in vivo</u> and <u>in vitro</u> tests are employed with the cytogenetic and microbial test systems. These tests and their descriptions are referenced in the Appendices A through F.

C. Compound

Test Material

Compound FDA 71-36, Calcium Propionate, Pfizer sample number 73211, as supplied by the Food and Drug Administration.

2. Dosages

The animals employed, the determination of the dosage levels and the route of administration are contained in the technical discussion.

The dosage levels employed for compound FDA 71-36 are as follows for the Cytogenetic Studies $\underline{\text{in vivo}}$ in rats.

Low Level	50 mg/kg
Intermediate Level	500 mg/kg
LD ₅	5000 mg/kg
Negative Control	Saline
Positive Control (TEM*)	0.3 mg/kg

The dosage levels employed for compound FDA 71-36 are as follows for the Host-Mediated Assay <u>in vivo</u> in mice.

Low Level	50 mg/kg
Intermediate Level	500 mg/kg
LD5	5000 mg/kg
Negative Control	Saline
Positive Control (EMS**)	350 mg/kg
(DMN***)	100 mg/kg

* Triethylene Melamine** Ethyl Methane Sulfonate*** Dimethyl Nitrosamine



The dosage levels employed for compound FDA 71-36 are as follows for the Dominant Lethal Assay $\underline{\text{in vivo}}$ in rats.

Low Level	50 mg/kg
Intermediate Level	500 mg/kg
LD ₅	5000 mg/kg
Negative Control	Saline
Positive Control (TEM*)	0.3 mg/kg

The <u>in vitro</u> Cytogenetic Studies were performed employing three logarithmic dose levels.

Low Level	0.4 mcg/m
Medium Level	4.0 mcg/ml
High Level	40.0 mcg/ml
Negative Control	Saline
Positive Control (TEM*)	0.1 mcg/ml

^{*}Triethylene Melamine

The discussion of this test is contained in the technical discussion.

D. <u>Methods</u>

The protocols employed are explained in Appendices C and D.

E. <u>Summary</u>

Host-Mediated Assay

Compound FDA 71-36 produced significant increases in the reversion frequency of <u>Salmonella</u> strain G-46 in the subacute tests. The increases were unrelated to dose levels. Lower, but similar responses were obtained at the two highest acute doses with G-46.

Strain TA-1530 showed no significant increases in reversion at either the acute or subacute doses.

A single dose was marginally recombinogenic in the acute trials using <u>Saccharomyces</u> strain D3. None of the other acute or subacute doses were recombinogenic against this organism.

All in vitro tests were negative.



2. Cytogenetics

a. <u>In vivo</u>

The compound produced no detectable significant aberration of the bone marrow metaphase chromosomes of rats when administered orally at the dosage levels employed in this study.

b. <u>In vitro</u>

The compound produced no significant aberration in the anaphase chromosomes of human tissue culture cells when tested at the dosage levels employed in this study.

3. Dominant Lethal

This compound was considered to be non-mutagenic in rats in the Dominant Lethal Assay when using the dosages employed in this study.

F. Results and Discussion

1. Toxicity Data

a. <u>In vivo</u>

Two male rats with an average body weight of 380 grams were given compound FDA 71-36 on an acute basis of 5000 mg/kg of body weight. The compound was in a solution of 0.85% saline and was administered by gastric intubation. The animals appeared normal during treatment and for an additional nine days post-treatment observation. Necropsies of these animals on day ten revealed no gross morphological changes in the organs examined. The work was repeated with a group of ten male albino rats with an average body weight of 380 grams with the same findings. In the experiment, 5000 mg/kg was administered at the high level, 500 mg/kg at the intermediate level, and 50 mg/kg at the low level. Animals in the acute studies were given a single dose

of the compound. The subacute study animals were given the same dosages as those in the acute study each day for five consecutive days, 24 hours apart.

b. <u>In vitro</u>

The compound was suspended in 0.85% saline at the concentrations listed above. It was introduced into culture tubes containing WI-38 cells in a logarithmic phase of growth. The cells were observed for cytopathic effect (CPE) and the presence of mitosis at 24 and 48 hours.

Tube No.	No. of Cells	Conc. mcg/ml	<u>CPE</u>	<u>Mitosis</u>
1	5 x 10 ⁵	1000	+	=
2	5 x 10 ⁵	1000	+	-
3	5 x 10 ⁵	500	+	-
4	5 x 10 ⁵	500	+	+
5	5 x 10 ⁵	100	+	
6	5 x 10 ⁵	100	+	+
7	5 x 10 ⁵	50	-	-
8	5 x 10 ⁵	50		+
9	5 x 10 ⁵	10	-	+
10	5 x 10 ⁵	10	-	+

Since both a presence of CPE and mitotic inhibition were observed, a closer range of concentrations was employed as follows.

Tube No.	No. of Cells	Conc. mcg/ml	<u>CPE</u>	<u>Mitosis</u>
1	5 x 10 ⁵	100	+	+
2	5 x 10 ⁵	. 100	-	+
3	5 x 10 ⁵	80	-	+
4	5 x 10 ⁵	80	+	+
5	5 x 10 ⁵	60	<u>+</u>	+
6	5 x 10 ⁵	60	-	+
7	5 x 10 ⁵	40	-	+
8	5 x 10 ⁵	40	-	+
9	5 x 10 ⁵	20	-	+ ,
10	5 x 10 ⁵	20	- ,	+

The high level was determined to be 40 mcg/ml since this level was free from CPE and mitotic inhibition. The intermediate level was 4.0~mcg/ml and the low level was 0.4~mcg/ml.



C. TOXICITY DATA SHEETS

CONTRACT FDA 71-268

COMPOUND FDA 71-36

CALCIUM PROPIONATE



TOXICITY DATA

COMPOUND FDA 71-36

Solvent:

0.85% saline

Dosage Form:

Suspension

Animals:

Male rats with an average body weight of 380 grams. All

animals were observed for ten (10) days.

Range Finding:

Dose mg/kg	No. Dead/No. Animals	Necropsy and Day of Death
5000	0/2	None
5000	0/10	None .

There were no abnormal gross pathology findings in the animals dosed at 5000 mg/kg and a determination of a $\rm LD_{50}$ was not performed.



2. Host-Mediated Assay

Compound FDA 71-36 produced significant increases in the reversion frequency of <u>Salmonella</u> strain G-46 in the subacute tests. The increases were unrelated to dose levels. Lower, but similar responses were obtained at the two highest acute doses with G-46.

Strain TA-1530 showed no significant increases in reversion at either acute or subacute doses.

A single dose was marginally recombinogenic in the acute trials using <u>Saccharomyces</u> strain D3. None of the other acute or subacute doses were recombinogenic against this organism.

Repeat tests of the subacute trials indicated the compound induced reversion, although the results were dose-independent.

All in vitro tests were negative.



Compound: 71-36 Calcium Propionate

			n Vivo	
Indicator Strain	<u>In Vitro</u>	Possible Low Recoveries	Controls	Other Comments
TA-1530	pos.	NC PC	NC OK	1. In cases where the recov. are somewhat low, the
11/1/72 PC NCSM Subacotes	neg.)	AL AI	PC OK	frequencies appear to be slightly elevated but
Subacoles		(AH) (SANC)	SANC OK	are generally in line with what is expected. Shouldn't
1/3/72 NC Acutes	•	SAL SAI SAH		present any problem.
		ЭМП		
	• .			
G-46				1. Acute doses show
		NC	NC OK	increase and respond
5/19/72	pos.	PC AL	PC A littl	to dose level
	neg.	AI	TO A TICE!	2. Subacute doses show
		AH	SANC OK	increase
	e.	SANC		
•		(SAL) SAI		
		SAH	•	
D 3				1. In the acute low dose
5/1/72 noutes	200	NC BC	NC OK	animal #2 does appear to
	pos.	PC AL	PC OK	be an outlier and probably should not be
5/5/72 Subacutes	(neg.)	AI		considered in calculati
•		HA	SANC OK	final freq.
•		SANC		2. The acute high dose loo
		SAL		as though it is positi
		SAI		All other dose levels a

Summary:

I feel that the data is acceptable. Historically the G-46 positive control is higher than the TA-1530 positive control. The data for G-46 appears to be consistent so that the lower than expected positive control is not a result of outliers or an unusually high negative control. G-46 results indicate that this compound is mutagenic in vivo. I also feel that the D3 acute high dose is positive.

I would expect this report to be accepted.

negative.

SAH

Compound: FDA_71_36 (Repeat)

	•		In Vivo	
Indicator Strain I	n Vitro	Possible Low Recoveries		Other Comments
TA-1530	pos.	NC PC	NC	
		AL	PC	
	neg.	AI		•
	-	НА	SANC	•
•		SANC		
Not repeated		SAL		
-		SAI		
,		SAH		
1				
- 47	•			
G-46		410	··~ OK	1. All three doses appear
Subacutes. Only		NC DC	NC OK	positive even though
Dunacu cos. Villy	pos.	PC ³	0.0 × 24	no dose response is evadent.
		AL. AI	PC OK	
	neg.	AH A1	SANC OK	
		SANC	SANC OR	
		SAL		
		SAI		
		SAH		
•			•	
D3		- -		
		NC	NC	
	pos.	PC		
		AL	PC	
Not repeated	neg.	AI	04110	
		AH	SANC	
		O A LI O		
		SANC		
		SAL	•	

Summary: This compound shows mutagenic activity in G-46 in the subacute doses when compared to the positive and negative controls. The lack of a good dose response is somewhat troublesome although many positive compounds fail to show dose responses in this system. I feel these data should be accepted.

a. HOST-MEDIATED ASSAY SUMMARY SHEETS

CONTRACT FDA 71-268

COMPOUND FDA 71-36

CALCIUM PROPIONATE



HOST MEDIATED ASSAY. SUMMARY SHEET.

COMBONIND:	FDA	•	SALMONELLA TA1530 G-4				YCES D-3	
		MMF (X 10E-8)	MFT/MFC.	MMF (X 10E-8)	MFT/MFG	MRF (X 10E-5)	MRT/MRC	
ACUTE, NC PC, AL, AI, AH,		.58 8.00 .75 .66	13.79 1.29 1.14 3.07	.54 9.51 .70 2.85 3.08	17.61 1.30 5.28 5,70	4,80 56,96 9,32 7,72 17,82	11.87 1.94 1.61 3.71	
SUBACUTE NC, SL, SI, SH	•	.83 1.48 1.43 1.14	1.78 1.72 1.37	.54 8.31 9.88 6.15	15.39 18.30 11.39	4,63 4,89 4,35 5,37	1.06 .94 1.16	
IN VITRO		TA1530 - - +	6 -46 - - +	% CONC 0.5 - 1.0	D-3 % SURVIVAL 71.8 100.0 42.7	R X 10E 7 5 381	5	

HOST MEDIATED ASSAY (REPEAT)

SUMMARY SHEET

COMPO	UND:	FDA	71-36

COMPOUND:	FDA (T-3)	5	SALMO	NELLA		SACCHAROMY	CES D-3
		TA153	30	G-46	•		
· • ···	(x	MMF 10E-8)	MFT/MFC	MMF (X 10E-8)	MFT/MFC	MRF (X 10E-5)	MRT/MRC
ACUTE							
NC	•	1.00	. •••	.67		1.00	_
PC		0.	0.	14.65	21.87	0.	0.
AL	<u>;</u> (0.	0.	0.	0.	0.	0.
AI).	0.	0.	0.	0.	0.
АН	. () .	0.	0.	0.	0.	0.
SUBACUTE	·						
NC		1.00		.67		1.00	
SL		0.	0.	5.20	7.76	0.	0.
SI		.	0.	7.21	10.76	0.	0.
SH		D. •	0.	4.60	6.87	0.	0.
IN VITRO		TA1530	G-46		D-3		•
				% CONC	% SURVIVA	L R X 10E	5

NC

PC

STOP SRU'S:.6

ISWITCH INS:MC602

b. HOST-MEDIATED ASSAY DATA SHEETS

CONTRACT FDA 71-268

COMPOUND FDA 71-36

CALCIUM PROPIONATE

Host Mediated Assay - Adjusted Faw CFU x 107/0.6 ml

Step 1: Technician set counter - plates on counter.

Step 2: Automatic equipment accumulates counts on 3 plates of 10⁻⁶ dilution as CFU x 10⁷/0.6 ml.

Step 3: Automatic equipment multiplies count obtained in step 1 by 0.1666666666667 to obtain total count/ml at 108.

Step 4: Automatic check of result of step 3. TC x $10^8 \div 0.1667 = \text{CFU x } 10^7/0.6 \text{ ml}$

Step 5: Technician was to record the true raw CFU x 107/0.6 ml in log book, however, the computer developed a quirk and provided the Column B check figure as the raw count.

To clarify the problem Column A is headed Adjusted Raw CFU X 10E⁷/0.6 ml in each case where the check figure was provided as the raw count.

COMPOUND: FUA 71-36

ORGANISM: SALMONELLA TA1530

DOSE LEVEL: NEGATIVE CONTROL - SALINE

TREATMENT: IN VIVO. ORAL. ACUTE

DATE STARTED: NOVEMBER 3, 1972

		A	В	С	D
	ANIMAL NUMBER	RAW CFU X 10E7/0.6ML	TOTAL CFU X 10E6/1.0ML	TOTAL NO. MUTANTS X 10E0/1.0ML	HUTATION FRE (C/B) X 10E-5
C	1	31.70	5.28	3.00	•57
	2	35.20	5.87	2.00	. 54
		21.70	3.62	2.00	•55
	· , 4	47.00	7.83	1.60	•13
	5	27.10	4.52	4.00	•69
*	: 6	17.60	2.93	3.00	1.02
	7	. 22.50	3.75	2.00	•53
	8	34.00	5.67	3.00	• 53
ц	9	43.80	7.30	5.00 -	•68
	Sin nor su			•	

NO. OF ANIMALS EQUALS 9
-SAMPLES WITH ZERO MUTANTS EQUAL 1

		COL. B	COL. C (X 10E0)	COL. D (X 101-8)
			-	
-	MEAN	5.20	2.78	•58
	RANGE	4.90	4.00	. 90
reference and the second second	MAX	7.83	5.00	1.02
	MIN .	2.93	1.00	.13

NO OUTLIERS

(X 10/8) (X 10E0) (X 10F-8 MEAN 3.67 2.75 .75 KANGE 4.63 6.06 1.10 MAX 6.30 7.00 1.41	DOSE LEGEL	: LOW - 50 MG/	/KG		
ANIMAL RAW CFU X. TOTAL CFU X MUTANTS X FRE (C/R NUMBER 10E7/0.6ML 10E8/1.0ML 10E0/1.0ML X 10E-0 1 29.70 4.95 7.00 1.41 2 37.80 6.30 2.00 .32 3 13.50 2.25 3.00 1.33 4 30.70 5.12 3.00 .59 5 24.10 4.02 2.00 .59 6 20.70 3.45 2.00 .50 7 8.80 1.47 1.60 .63 7 8.80 1.47 1.60 .63 8 20.50 3.42 2.00 .59 NO. OF ANIMALS EGUALS 8 TOTAL CFU OUT OF RANGE EGUALS 2 COL. B COL. C COL. D (X 10E8) (X 10E0) (X 10F-8 RANGE 4.85 6.05 1.10 MAX 6.30 7.00 1.41 MIN 1.47 1.00 .32	TREATMENT:	IN VIVO. GRAU	. ACUTE	DATE STARTED:	NOVEMBER 3
ANIMAL RAW CFU X. TOTAL CFU X MUTANTS X FRE (C/R. NUMBER 10E7/0.6ML 10E8/1.0ML 10E0/1.0ML X 10E-0 1 29.70 4.95 7.00 1.41 2 37.30 6.50 2.60 .32 3 13.50 2.25 3.00 1.33 4 30.70 5.12 3.00 .50 5 24.10 4.02 2.00 .50 6 20.70 3.45 2.00 .50 7 8.80 1.47 1.00 .63 7 8.80 1.47 1.00 .63 5 20.50 3.42 2.00 .50 NO. OF ANIMALS EQUALS 8 TOTAL CFU OUT OF RANGE EQUALS 2 COL. B COL. C COL. D (X.10L8) (X.10E0) (X.10F-8 NANGE 4.83 6.50 7.00 1.41 MAX 6.30 7.00 1.41 MIN 1.47 1.00 .32	The section of the se	Α	В		
NUMBER 10E7/0.6ML 10E8/1.0ML 10E0/1.0ML X 10E-0 1 29.70 4.95 7.00 1.41 2 37.80 6.50 2.00 .32 3 13.50 2.25 3.00 1.33 4 30.70 5.12 3.00 .59 5 24.10 4.02 2.00 .50 6 20.70 3.45 2.00 .58 7 8.80 1.47 1.00 .63 8 20.50 3.42 2.00 .59 NO. OF ANIMALS EQUALS 8 TOTAL CEU OUT OF RANGE EQUALS 2 COL. B COL. C COL. D (X 10E8) (X 10E6) (X 10F-8 RANGE 4.63 6.00 1.10 MAX 6.30 7.00 1.41 MIN 1.47 1.00 .32					
1 29.70 4.95 7.00 1.41 2 37.30 6.50 2.60 .32 3 13.50 2.25 3.00 1.33 4 30.70 5.12 3.00 .50 5 24.10 4.02 2.00 .50 6 20.70 3.45 2.00 .50 7 8.80 1.47 1.00 .63 8 20.50 3.42 2.00 .50 NO. OF ANIMALS EQUALS 8 TOTAL CFU OUT OF RANGE EQUALS 2 COL. B					
2 37.80 6.50 2.00 .32 3 13.50 2.25 3.00 1.33 4 30.70 5.12 3.00 .59 5 24.10 4.02 2.00 .50 6 20.70 3.45 2.00 .53 7 8.80 1.47 1.00 .63 6 20.50 3.42 2.00 .59 NO. OF ANIMALS EQUALS 8 TOTAL CFU OUT OF RANGE EQUALS 2 (X 1028) (X 1060) (X 107-8 MEAN 3.67 2.75 .79 RANGE 4.63 6.00 7.00 1.41 MIN 1.47 1.00 .32	MOSSEK	TOCIVATIONE	TOUGNION	TOUNT	X 10ETC
2 37.80 6.50 2.00 .32 3 13.50 2.25 3.00 1.33 4 30.70 5.12 3.00 .59 5 24.10 4.02 2.00 .50 6 20.70 3.45 2.00 .53 7 8.80 1.47 1.00 .63 6 20.50 3.42 2.00 .59 NO. OF ANIMALS EQUALS 8 TOTAL CFU OUT OF RANGE EQUALS 2 (X 1028) (X 1060) (X 107-8 MEAN 3.67 2.75 .79 RANGE 4.63 6.00 7.00 1.41 MIN 1.47 1.00 .32	i	29.70	4.95	7.00	1.41
3 13.50 2.25 3.00 1.33 4 30.70 5.12 3.00 .59 5 24.10 4.02 2.00 .50 6 20.70 3.45 2.00 .58 7 8.80 1.47 1.00 .63 6 20.50 3.42 2.00 .50 NO. OF ANIMALS EQUALS 8 TOTAL CFU DUT OF RANGE EQUALS 2 COL. B COL. C COL. D (X 10.8) (X 10E0) (X 10F-8 RANGE 4.83 6.00 1.10 MAX 6.30 7.00 1.41 MIN 1.47 1.00 .32					
5 24.10 4.02 2.00 .50 5 20.70 3.45 2.00 .58 7 8.80 1.47 1.00 .63 6 20.50 3.42 2.00 .50 NO. OF ANIMALS EQUALS 8 TOTAL CFU DUT OF RANGE EQUALS 2 COL. B COL. C COL. D (X 10/8) (X 10E6) (X 10F-8 MEAN 3.67 2.75 .75 RANGE 4.63 6.00 7.00 1.41 MIN 1.47 1.00 .32	3				
5 20.70 3.45 2.00 .58 7 8.80 1.47 1.00 .63 6 20.50 3.42 2.00 .59 NO. OF ANIMALS EQUALS 8 TOTAL CFU OUT OF RANGE EQUALS 2 COL. B COL. C COL. D (X 10:8) (X 10:6) (X 10:6) MEAN 3.07 2.75 .75 KANSE 4.83 6.00 7.00 1.41 MAX 6.30 7.00 1.41 MIN 1.47 1.00 .32			5.12	3.00	
7 8.80 1.47 1.00 .63 6 20.50 3.42 2.00 .50 NO. OF ANIMALS EQUALS 8 TOTAL CFU DUT OF RANGE EQUALS 2 COL. B COL. C COL. D (X 10[8) (X 10E0) (X 10F-8 MEAN 3.67 2.75 .75 RANGE 4.83 6.00 1.10 MAX 6.30 7.00 1.41 MIN 1.47 1.00 .32					
NO. OF ANIMALS EQUALS 8 TOTAL CFU DUT OF RANGE EQUALS 2.00 .50					
NO. OF ANIMALS EQUALS 8 TOTAL CFU OUT OF RANGE EQUALS 2 COL. B	7				
TOTAL CFU DUT OF RANGE EQUALS 2 COL. B COL. C COL. D (X 10[8]) (X 10E0) (X 10F-8 MEAN		20.50		2.00	
(X 10/8) (X 10E0) (X 10F-8 MEAN 3.67 2.75 .75 KANGE 4.83 6.05 1.10 MAX 6.30 7.00 1.41 MIN 1.47 1.00 .32	NO. OF ANI TOTAL CEU	MALS EQUALS DUT OF RANGE (EGUALS 2		
MEAN 3.67 2.75 .75 RANGE 4.83 6.00 1.10 MAX 6.30 7.09 1.41 MIN 1.47 1.00 .32					col. D
RANGE 4.83 6.05 1.10 MAX 6.30 7.03 1.41 MIN 1.47 1.00 .32	enterente de tente e pro- la relación de la relació	EAC AN			
MAX 6.30 7.00 1.41 MIN 1.47 1.00 .32	.•				
MIN 1.47 1.00 .32		and the second s			
NO OUTLIERS	THE STATE OF THE S				.32
	NO OUTLIER	5			
		management of the following states of the st	Andrew paper of the control of the c	CONTRACTOR OF THE MINISTER SAME ASSESSMENT OF MINISTER ABOVE CAMBER OF THE	

COMPOUND: FOA 71-36	ORGANISM:	SALMONELLA	TA1530
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DOSE LEVEL: INTERMEDIATE - 500 MG/KG

٧.) (تعا	TREATMENT	: IN VIVO. ORAL	. ACUTE	DATE STARTED:	NOVEMBER 3.	197
		A	В	C TOTAL NO.	D	
	ANIMAL NUMBER	RAW CFU X 10E7/0.6ML	TOTAL CFU X 10EU/1.0ML	MUTANTS X 10E0/1.0ML	FRE (C/B) X 10E-8	• • "
	1	12.70	2.12	2.00	• 94	
Lt.	2	71.20	11.87	7.00	•59	
		···· 61•60	10.27	8,00	· - 73 · · ·	
	' 4	46.00	7.67	4.00	•52	
U :	5	9.20	1.53	1.60	• 6 5	
		47.20	7•87	5.00	•64	
	7	3+80	1.47	1.00	•68	
	8	40.20	6.70	3.00	•45	
·		21.40	3.57	3.00	•64	
<u></u>	10	32.70	5.45	3.00	•55	•
	NO. OF AN	IMALS EQUALS	10		• ,	
	-		COL. B	COL. C	COL. D	
U		The second secon	(X 10E8)	(X 10E0)	(X 10世-8)	
- i. ,		MEAN	5.85	3.70	• 56	
		RANGE	10.40	7.00	•50	
 	e de la companya de l La companya de la co	MAX	11.87	5.00	• 94	
ч.	A4	MIN	1.47	1.00	•45	
Film.	NO OUTLIE	R5				

COMPOUND: FDA 71-36

ORGANISM: SALMONELLA TA1530

DOSE LEVEL: HIGH - 5000 MG/Kg

TREATMENT: IN VIVO. ORAL, ACUTE

DATE STARTED: NOVEMBER 3, 1972

	Α	8	C	ס
ANIMAL NUMBER	RAW CFU X 10E7/0.6ML	TOTAL CFU X 10LB/1.0ML	MUTANTS X 10E0/1.0HL	HUTATION FRE (C/B) X 10E-R
1	9.80	1.63	2.00	1.22
2	9.40	1.57	3.00	1.91
3	11.80	1.97	4.00	2.03
4	12.00	2.00	4.00	2.00
5	14.60	2.43	5.00	2.05
6	9.20	1.53	3•00	1.95
7	18.20	3.03	4.00	1.32
8	16.90	2.62	5+00	1.78

NO. OF ANIMALS EQUALS 8
TOTAL CFU OUT OF RANGE LOUALS

	COL. B	COL. C	COL. D
•	(X 10E8)	(X 10Eg)	(X 10E-8)
MEAN	2.12	3.75	1.78
RANGE	1.50	3.00	•83
MAX	3.63	5.00	2.05
MIN	1.53	2.00	1.22

NO OUTLIERS

COMPOUND: FDA 71-36

ORGANISM: SALMONELLA TA1530

DOSE LEVEL: NEGATIVE CONTROL - SUBACUTE TRIALS

TREATMENT: IN VIVO, ORAL, ACUTE

DATE STARTED: NOVEMBER 1, 1972

		A	В	C TOTAL NO	D MUTATION
	ANIMAL NUMBER	RAW CFU X 10E7/0.6ML	TOTAL CFU X 10E8/1.0HL	HUTANTS X 10EO/1.0ML	FRE (C/B)
	1	8∙70	1.45	3.00	2.07
	2	10.40	2.73	2.00	.73
: 			2.97	1.00	
	. 4	16.80	2.60	1.00	• j
	5	12.80	2.13	1.00	•47
	6	13.40	2.23	2.00	•90
	7	16.20	2.70	2.00	.74
3	8	22.70	3.78	4.00	1.06

NO. OF ANIMALS EQUALS 8
TOTAL CFU OUT OF RANGE EQUALS 1
SAMPLES WITH ZERO MUTANTS EQUAL 1

	COL. B	COL. C	COL. D
er er er caminicación en	(X 10E8)	(X 10E0)	(X 10E-8)
MEAN	2.60	2.00	-83
RANGE	2.33	3.00	1.73
MAX	3.78	4.00	2.07
MIN	1.45	1.00	•34

* SUMMARY WITH OUTLIERS REMOVED

			MEAN	(X	10E8)	· (X)L. C 10E0)	
ľ			RANGE MAX		1.65 3.78	e en en en egy en ennem .	3.00 4.00	•66 •72 1•06
	en e	i de la companya de La companya de la co	MIN		2.13		1.00	34

X CSC85F 04 DEC 72 13:44:18 USER CFU007 200.

DS IN 346 OUT O LINES 276 PROCESSING TIME 13.

13. 9 SECONDS

RNATE PRINT FILE - RUN 200 CFU007, NUMBER= 200

PAP+6 PRINT ON SIX-PART PAPER

COMPOUND: FDA 71-36

ORGANISM: SALMONELLA TA1530

DOSE LEVEL: POSITIVE CONTROL - DAN - 100 MG/KG

TREATMENT: IN VIVO. ORAL, ACUTE - DATE STARTED: NOVEMBER 1, 1972

		., B	C	D
ANIMAL NUMBER	ADJUSTED RAW CFU X 10E7/0.6ML	TOTAL CFU X 10E8/1.0ML	TOTAL NO. MUTANTS X 10E0/1.0ML	MUTATION FRE (C/B) X 102-6
1	35+20	5.87	38.00	6.48
2	22.50	3.80	42.00	11.05
3	31.70	5.28	28.00	5.30
4	19.20	3.20	30.00	9.37
		4.73	22.00	4.05
б	31.00	5.17	36.00	6.97
7	23.10	3.85	47.00	12.21

NO. OF ANIMALS EQUALS NO. OF CONTAMINATED EQUALS - TOTAL CFU OUT OF RANGE EQUALS

	COL. B	COL. C	COL. D
en e	(X 10E8)	(X 10E0)	- (X 10E-8)
MEAN	4.56	34.71	ë•00
RANGE	2.67	25.00	7.56
MAX	5.87	47.00	12.21
MIN	3.20	22.00	4.65

K CSC85F 04 DEC 72 13:45:17 USER CFU007 200

75 PROCESSING TIME LINES

COMPOUND: FDA 71-36 ORGANISM: SALMONELLA TA1539

GOSE LEVEL: LOW - 50 MG/KG

TREATMENT: IN VIVO, OKAL, SUBACUTE DATE STARTED: NOVEMBER 1, 1972

	A	8	C	Ð
ANIMAL NUMBER	RAW CFU X 1067/0.6HL	TOTAL CFU X 1028/1.0ML	TOTAL NO. MUTANTS X 10E0/1.0ML	MUTATION FRE (C/B) X 10E-3
1	7.50	1.25	3.00	2.40
Ż	39.00	6.50	2.00	• 31
3	18.20	3.03	1.00	•33
ц	21.00	3.50	3.00	•86
5	11.40	1.90	2+90	1.05
6	11.80	1.97	2.00	1.02
7	6.80	1.13	5.00	4.41 .

NO. OF ANIMALS EQUALS 7
NO. OF CONTAMINATED EQUALS 1
TOTAL CFU OUT OF HANGE EQUALS 1
SAMPLES WITH ZERO MUTANTS EQUAL 1

	COL. B	COL. C	COL. D
	(X 160a)	· (X 1086)	(8-301 X)
MEAN	2.75	2.57	1.48
RANGE	5.37	4.00	4.10
XAM	6.50	5.00	4.41
MIN	1.13	1.00	.31

* SUMMARY WITH OUTLIERS REMOVED

	COL. B	COL. C	cot. D
	(x 1008)	(X 10E0)	(X 10E-8)
ME A.1	3.63	2.17	• 79
RANGE	5 • 25	2.00	2.09
MAX	6+50	3.00	2.40
MIN	1.25	1.00	

COMPOUND: FDA 71-36

ORGANISM: SALMONELLA TA1550

DOSE LEVEL: INTERMEDIATE - 500 MG/KG

TREATMENT: IN VIVO. CRAL, SUBACUTE

DATE STARTED: NOVEMBER 1. 1972

	A	В	· c	D
ARIMAL NUMBER	RAW CFU X 1067/0.6ML	TOTAL CFU X 1028/1.0ML	TOTAL NO. MUTANTS X 10E0/1.0ML	HUTATION FRE (C/B) X 10E-A
1 2	13.30 10.20	2.22 1.70	8.00 3.00	3.61
3 4	35.60 14.60	5.95 2.47	10.00 2.00	1.75 1.69
5	43.10	7.13	4.00	•31 •56
7	30+10 28∙30	5•02 4•78	5•00 6•00	1•00 1•27
.	31.00	5+17	4.00	. 77

NO. OF ANIMALS EQUALS 8
NO. OF DEAD ANIMALS EQUALS 1
TOTAL CFU OUT OF RANGE EQUALS 1

	COL. B	. COL. C	cor. D	
	(X 1008)	(X 10E0)	(X 105-8)	
MEAN	4.30	5.25	1.43	
RANGE	5•48	8.00	3.05	
MAX	7.18	10.00	3.61	
MIN	1.70	2.00	•56	

* SUMMARY WITH OUTLIERS REMOVED

$(\hat{\mathbf{x}}_{i}) = \frac{1}{2} \left(\hat{\mathbf{x}}_{i} \cdot \hat{\mathbf{x}}_{i} \right) = \hat{\mathbf{x}}_{i} \cdot \hat{\mathbf{x}}_{i} = 0$	COL. B	COL. C	COL. D
	(X 10c8)	(x 10E0)	(X 105-8)
MEAN	4.00	4.66	1.12
RANGE	5.48	8.00	1.21
MAX	7.18	10.0c	1.76
HIN	1.70	2.00	• 56

COMPOUND: FDA 71-36 ORGANISM: SALMONELLA TA1530 DOSE LEVEL: HIGH - 5000 MG/KG TREATMENT: IN VIVO, ORAL, SUBACUTE DATE STARTED: NOVEMBER 1, 1972 TOTAL NO. MUTATION ANIMAL RAW CFU X TOTAL CFU X MUTANTS X FRE (C/B) 10E8/1.UML NUMBER 10E7/J.6ML 1026/1.0ML X 10E-8 49.60 8.27 2.00 6.30 .95 1.05 1.00 3 1.23 7.40 2.00 1.62 16.00 2.67 2.00 •75 7.40 1.25 3.00 2.43 32.10 5.35 .75 4.00 24.00 4.00 5.00 1.25 NO. OF ANIMALS EQUALS TOTAL CFU OUT OF KANGE EQUALS COL. B COL. C COL. D (X 1088) (X 10E0) (X 105-8) MEAN 3.40 2.71 1.14 RANGE 7.22 4.00 2.19 MAX. 8.27 5.00 2.43 MIN 1.00 1.05 . 24 * SUMMARY WITH OUTLIERS REMOVED COL. i3 COL. C COL. D (X 1008) (X 10E8) (X 10E-8) HE AN 3.76 2.67 • 03 RANGE 7.22 4.00 1.38 MAX 6.27 5.00 1.62 MIN 1.00 1.05

15:20:49 USER CFUGG7

200

526 PROCESSING TIME

MEX CSCBSF 02 DEC 72

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19.65 SECONDS

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and the second of the second of the second	HOST ME	DIATED ASSAY R	EPORT SHEET	
	IND: FUA 71-36		ORGANISM: SAL	HONELLA G-46
DOSE L	EVEL: NEGATIVE CO	NTROL - SALINE		
TREATM	ENT: IN VIVO, ORA	L. ACUTE	DATE STARTED:	MAY 19, 1972
	A	8	C.	D
ANIMAL	ADJUSTED RAW CFU X	TOTAL CFU X	TOTAL NO.	
NUMBER		10E8/1.0ML	MUTANTS X 10EU/1.0ML	FRE (C/6) % 10E-8
1	26•26	4.71	1.00	•21
2	54.96	9.16	4.00	• 44
4	24 • 43 40 • 14	4.08	1.00	• 25
5	34.08	6•69 5•68	5.00	• 75
6	37.38	6•23	7.00 4.00	1.23
7	20.22	3.37	1.00	• 30
8	31.44	5.24	4.00	•76
9	60.72	10.12	3,00	······································
NO. OF	ANIMALS EQUALS	0	•	
TOTAL	CFU OUT OF RANGE	EQUALS 1		
		COL. B	COL. C	201 5
	-	(X 10E8)	(X 10E0)	COL. D
	MEAN	6.14	3.33	•54
	RANGE	6.75	5.00	1.02
	MAX	10.12	7.00	1.23
	MIN	3,37	1.00	.21
	er over the single control of the second	ere ej erekon ili erekon ili ili birak	in the second of	
	*	SUMMARY WITH (OUTLIERS REMOVE)
	•			
	e e e e e e e e e e e e e e e e e e e	COL. B	COL. C	COL. D
	e e e e e e e e e e e e e e e e e e e	(X 10E8)	(X 1058) COF* C	COL. D (X 101-8)
	MEAN	(X 10E8) 6.20	(0201 X) 88•s	
	MEAN RANGE MAX	(X 10E8)	(Y topo)	(X 10L-8)

COMPOUND: FDA 71-36

STOP

ORGANISM: SALMONELLA G-46

DOSE LEVEL: POSITIVE CONTROL - DMN - 100 MG/KG

TREATMENT: IN VIVO, ORAL, ACUTE

DATE STARTED: MAY 19: 1972

s ktoria s	A ADJUSTED	8	C TOTAL NO.	D MUTATION
ANIMAL, NUMBER	RAW CFU X 10E7/0.6ML	TOTAL CFU X	MUTANTS X 10E0/1.0ML	FRE (C/B) X 10E-8
1	11,82	1,97 4,98	20,00 34,00	10.15
23456789	29,88 23,94	3,99	38,00	6.83 9.52
4.	22,68	3,78	24,00	6.35
5	33.78	5,63	.44.00	7.62
5	33.90	5,65	75.00	13.27
Ŕ,	20,13 39,20	3,36 6,53	23.00 97.00	6.85 14.85
9	37.70	6,28	61,00	9.71
10	32,55	5,43	53,00	9.77
NO. OF AN	IIMALS EQUALS	10	•	
		COL. B (X 10E8)	COL, C (X 10E0)	COL. D (% 10E-8)
•	MEAN	4.76	46.90	9,51
	RANGE	4.56	77.00	8,50
	MAX	6.53	97.00	14.85
NO OUTLIE	MIN RS	1.97	20.00	6,35

h, d					
7	COMPOUND:	FDA 71-36		ORGANISM: SAL	MONELLA 6-46
j	BACK I FVE	L: LOW - 50 MG,	ANG.		
	DOSE FEAC	To Ech is 20 May	, NO	en e	· · · · · · · · · · · · · · · · · · ·
	TREATMENT	: IN VIVO. ORAL	. ACUTE	DATE STARTED:	MAY 19: 1972
			and the second s	angana sa mangan sa	n ann an h-rin againe - r-rin - ar ar lainn an - ar
		A	8.	C	מ
J		ADJUSTED		TOTAL NO.	MUTATION
	ANIHAL		TOTAL CFU X		FRE (C/8)
	NUMBER	10E7/0.6ML	10E8/1.0ML	10EU/1.0%L	X 10E-8
1		29.70	4.95 ·····	3.00	
79	2	7.50	1.25	1,00	.60
	3	40.44	6.74	5.00	.74
J		53.10	B.85	5.00	
_	5	58.32	9.72	10.00	1.03
Î	6	25.50	4.25	2.00	•47
J		45.96	7.66	6.00	•78
	8	59.16	9.86	8.00	•01
	9	32-40	5.40	2,00	•37
	10	49.98	8 ₄ 33	7.00	∵
2	NO. OF AN	MIMALS EQUALS	10		
		nggan ang anggang na ang ang ang ang ang	COL. B	COL. C	COL. D
			(X 10E8)	(X 10E0)	(X 10E-8)
	e on section in	ME AN	······································		70
		RAHGE	8.61	9.00	•66
:::: 7		MAX	9.86	10.00	1.03
M	ganta spaniga area de la cale de	HIN	1.25	- 1.00	
3	NO GUTLIE	ERS			•
					
7		and the second s	and the second s	·	
j					
					
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-	n man andametra constitute Alexandra de la	ting	<u></u>	Committee that the second of t	and the second of the second o
7				•	
1					
#	The state of the property of the state of th	and the second s	and the second of the second o	and a second control of the second control o	grade come and grade an above of the second control of the second

TREATHENT	: IN VIVO, ORAL	L. ACUTE	DATE STARTED	MAY 19, 19
	A	₿	С	
A	ADJUSTED		TOTAL NO.	D MILTATION
ANIHAL	RAW CFU X	TOTAL CFU X	MUTANTS X-	MUTATION
NUMBER	10E7/0.6ML	10E8/1.0ML	10E0/1.0ML	FRE (C/E
1	40.14		4.00	
2 3	18.24	3.04	19.00	•60
3	24 • 84	4.14	18.00	6.25
	18.54	3.09	3.00	4 • 35
5	23.46	3.91	6.00	.97
6	25.86	4.31	4.00	1.53
	14.18	2.36	7.00	•93
8	30•36	5.06	26.00	2.95
9	42.81	7.14	21.00	5.14 2.94
NO. OF AN NO. OF DE	IMALS EQUALS AD ANIMALS EQUA	9 LS 1	The state of the s	<u></u>
		COL. B	COL. C	COL. D
		(X 10E8)	(X 1020)	(X. 102-8
meriod of the second	MEAN	4.42	17.00	2.35
	RANGE	4.77	23.00	ຂ•ບວ ວົ•ບໍ5
	MAX.	7.14	26.00	. 6. 25
NA ALIMI TO	MIN	2.36 ·· -	3.00	
NO DUTLIE	35		* - * -	• 00

	FDA 71-36		ORGANISM: SAL	MONELLA 6-46
	L: HIGH - 5000	The second secon		te de marco de la compansión de la compa
TREATMENT	: IN VIVO. ORAL	. ACUTE	DATE STARTED:	MAY 19, 1972
en e	A	В	C	D
	ADJUSTED		TOTAL NO.	MUTATION
ANIMAL				FRE (C/B)
NUMBER	10E7/0.6ML	10E8/1.GML	10E0/1.0ML	X 10E-8
	30.36	5.06	23,00	4.55
2 3	30.66	5.11	18.00	3.52
	31.30	5.30	16.00	3.02
			12.00	2.06
5	29•58	4.93	14.00	2.84
. 6	26•26	4.38	13.00	2.97
7 8	43.98	7.33	15.00	2.05
. 0	28.32	4.72	17.00	3. 60
NO. OF AN	IMALS EQUALS	g		-1
	AD ANIMALS EQUA	LS 2		
······································		COL. B	·········· COL. C	COL. D
		(X 10E8)		
	MEAN	5.33		3.03
	RAMGE	2.95		2.50
	MAX	7.33	23.00	4.55
	MIN.	4.38	12.00	2.05
The state of the s				
		SUMMARY WITH	OUTLIERS REMOVE	ip
		COL. B	col. c	COL. D
	and the specimen pairs a descript particular to the street where the section was an experiment.		(X 10E0)	(X-10£-8)
	MEAN	5.37	15.00	2.37
	RANGE	2.95	6.00	1.56
	MAX	7.33	1:00	3.60
•	MIN	4.38	12.00	2.05
		. + 4	2000	**

	COMPOUND:	FDA 71-36		ORGANISM: SALI	MONELLA G-46
. F. 9	DOSE LEVEL	: LOW - 50 MG/	YKG		
	TREATMENT:	IN VIVO. ORAL	. SUBACUTE	DATE STARTED:	MAY 19, 1972
		A ADJUSTED	B :	C TOTAL NO.	D HOITATUM
n	ANIMAL NUMBER	- RAW CFU X 10E7/0.6ML	TOTAL CFU X	MUTANTS X	FRE (C/6) X 10E-8
	2 3	22.50 17.82 20.34 29.28	3.75 2.97 3.39	45.00 42.00	5.07 15.15 12.39
	5 6	25.25 25.10 19.32 19.98	4.88 4.35 3.22 3.33	45.00 32.00 39.00 13.00	9.22 7.36 12.11 3.90
	8 9	18.18 17.94	3.03 2.99	17.00 12.00	5.01 4.01
		MALS EQUALS OUT OF RANGE D	9 EQUALS 1	·	
			COL. B (X 10E8)	COL. C (X 10E0)	COL. D (X 10L-8)
Season Name of Street	te de la companya de	MEAN RANGE MAX	3.55 1.91 4.68	20.33 33.00 43.00	8.31 11.25 15.15
	NO OUTLIER	RS			3.90
		. i., <u>. i</u>	<u> </u>		

COMF	POUND: FUA 71-36		ORGANISM: SAL	MONELLA 6-46
Dose	E LEVEL: INTERMEDIAT	TE - 500 MG/KG		
TRE	ATMENT: IN VIVO, OR	AL. SUBACUTE	DATE STARTED	: HAY 19: 1972
ANII	A ADJUSTED MAL RAW CFU X	B TOTAL CFU X	C TOTAL NO. MUTANTS X	D MUTATION FRE (C/8)
NUME			10E0/1.0ML	X 10E-8
1	43.98 29.94	7.33 4.99	70.00	2.57
2 3	29.94 34.66 21.30	5•78 3•55	51.60 60.00	14.03 8.82
5 6	28•68 23•74	4•78 4•79	55.00 42.00	16.90 11.51 8.77
8 9	-	5.83 2.63	61.00 48.00	9.08 8.23
10	26.52	4.42	36.00 33.00	11.41
No.	OF ANIMALS EQUALS	10		The second secon
	tem 63.5	COL. B (X 10E6)	COL. C (X 10E0)	COL. D (X 105-8)
	MEAN RANGE MAX	5.08 4.70 7.33	51.00 70.00	9.88 14.31 . 16.90
	MIN	2.63	19.00	2.59
		* SUMMARY WITH	OUTLIERS REMOV	ED
	MEAN	COL. B (X 10E6) 4.83	COL. C (X 10E0) 50.00	COL. D (X 106-8) 10.69
······································	RANGE MAX MIN	4.09 6.72 2.63	40.00 76.00 30.00	9.44 16.90 7.47

	TREATMENT	: IN VIVO, ORAL	, SUBACUTE	DATE STARTED:	MAY 19: 1972
	ANIMAL HUMBER	A AJUSTED PAW CFU X 10E7/0.6ML	B TOTAL CFU X 10E5/1.0ML	C TOTAL NO. MUTANTS X 10E0/1.0ML	D MUTATION FRE (C/b) X 10t-8
A	2	45.96 15.48 50.94	7.66 2.58 3.49	34.00 29.00 34.00	4.44 11.24 4.00
		29.94 30.66 32.52	4.99 5.11 5.42	21.00 32.00 13.00	4.21 6.20 3.32
	8 9	37.98 23.88 25.34	6.33 3.98 4.39	29.00 20.00 28.00	4.58 7.64 6.38
	10 • OF AN	19.08	3.18	32.00	. 18.00
		MEAN RANGE MAX MIN	COL. B (X 10E8) 5.21 5.91 6.49 2.58	COL. C (X 10E0) 23.50 16.00 34.00	COL. D (X 10E-8) 6.15 7.92 11.24
). 	NO OUTLI	•		15.00	Q. J.

RDS-IN-- 532 OUT --- 0 LINES -- 617 PROCESSING TIME -- 22.95 SECONDS

HOST MEDIATED ASSAY REPORT SHEET (REPEAT)

COMPOUND: FDA 71-36 ORGANISM: SALMONELLA G-46

DOSE LEVEL: NEGATIVE CONTROL - SALINE

TREATMENT: IN VIVO, ORAL, SUBACUTE DATE STARTED: JUNE 13, 1973

ANIMAL NUMBER	A RAW CFU X 10E7/0.6ML	B TOTAL CFU X 10E8/1.0ML	C TOTAL NO. MUTANTS X 10E0/1.0ML	D MUTATION FRE (C/B) X 10E-8
1 2 3 4 5 6 7 8 9	65.70 111.80 116.00 96.30 69.60 70.60 68.30 93.60 95.50 58.90	10.95 18.63 19.33 16.05 11.60 11.77 11.38 15.60 15.92 9.82	4.00 19.00 9.00 12.00 7.00 9.00 8.00 13.00 8.00 7.00	.37 1.02 .47 .75 .60 .76 .70 .83 .50

NO. OF ANIMALS EQUALS 10

		COL. B (X 10E8)	COL. C (X 10E0)	COL. D (X 10E-8)
	MEAN	14.11	9.60	.67
	RANGE	9.52	15.00	.65
	MAX	19.33	19.00	1.02
	MIN	9.82	4.00	• 37

NO OUTLIERS

STOP SRU'S:.7

HOST MEDIATED ASSAY REPORT SHEET (REPEAT)

COMPOUND: FDA 71-36 ORGANISM: SALMONELLA G-46

DOSE LEVEL: POSITIVE CONTROL - DMN - 100 MG/KG

TREATMENT: IN VIVO, ORAL, ACUTE DATE STARTED: JUNE 13, 1973

	Α	В	C TOTAL NO.	D MUTATION
ANIMAL	RAW CFU X	TOTAL CFU X	MUTANTS X	FRE (C/B)
NUMBER	10E7/0.6ML	10E8/1.0ML	10E0/1.0ML	X 10E-8
1	43.40	7.23	172.00	23.78
2	66.00	11.00	167.00	15.18
3	52.00	8.67	183.00	21.11
4	52.60	8.77	89.00	10.15
5	46.70	7.78	141.00	18.12
6	36.10	6.02	69.00	11.47
7	55.00	9.17	93.00	10.15
8	65.20	10.87	141.00	12.98
9	73.10	12.18	196.00	16.09
10	82.70	13.78	103.00	7.47

NO. OF ANIMALS EQUALS 10

	COL. B	COL. C	COL. D
	(X 10E8)	(X 10E0)	(X 10E-8)
MEAN	9.55	135.40	14.65
RANGE	7.77	127.00	16.31
MAX	13.78	196.00	23.78
MIN	6.02 .	69.00	7.47

NO OUTLIERS

STOP SRU'S:.6

!switch\$: INMC653

GPS144 E XQT PROG NAME > 6 CHAR ISWITHC INS:MC653

!SWITHC INS:MC653 SWITCH INS:MC653

!SAL

GT3104 E UNDER INED COMMAND

HOST MEDIATED ASSAY REPORT SHEET (REPEAT)

COMPOUND: FDA 71-36 ORGANISM: SALMONELLA G-46

DOSE LEVEL: LOW - 50 MG/KG

TREATMENT: IN VIVO, ORAL, SUBACUTE DATE STARTED: JUNE 13, 1973

ANIMAL NUMBER	A RAW CFU X 10E7/0.6ML	B TOTAL CFU X 10E8/1.0ML	C TOTAL NO. MUTANTS X 10E0/1.0ML	D MUTATION FRE (C/B) X 10E-8
1 2 3 4 5 6 7 8	50.70 43.70 42.30 45.90 76.60 57.50 47.50	8.45 7.28 7.05 7.65 12.77 9.58 7.92 17.28	23.00 49.00 24.00 24.00 64.00 77.00 44.00 121.00	2.72 6.73 3.40 3.14 5.01 8.03 5.56 7.00

NO. OF ANIMALS EQUALS 8
NO. OF CONTAMINATED EQUALS 2

	COL. B	COL. C	COL. D
	(X 10E8)	(X 10E0)	(X 10E-8)
MEAN	9.75	53.25	5.20
RANGE	10.23	98.00	5.31
MAX	17.28	121.00	8.03
MIN	7.05	23.00	2.72

NO OUTLIERS

STOP SRU'S:.6 !SWITCH IND:MC654 !SAL

HOST MEDIATED ASSAY REPORT SHEET (REPEAT)

COMPOUND: FDA 71-36 ORGANISM: SALMONELLA G-46

DOSE LEVEL: INTERMEDIATE - 500 MG/KG

TREATMENT: IN VIVO, ORAL, SUBACUTE DATE STARTED: JUNE 13, 1973

	··· A	В	C TOTAL NO.	D MUTATION
ANIMAL	RAW CFU X	TOTAL CFU X	MUTANTS X	FRE (C/B)
NUMBER	10E7/0.6ML	10E8/1.0ML	10E0/1.0ML	X 10E-8
1	128.50	21.42	142.00	6.63
2	83.80	13.97	64.00	4.58
3	65.60	10.93	72.00	6.59
4	113.90	18.98	114.00	6.01
5	57.30	9.55	84.00	8.80
6	53.50	8.92	82.00	9.20
7 .	58.20	9.70	72.00	7.42
8	81.10	13.52	94.00	6.95
9	50.80	8.47	74.00	8.74

NO. OF ANIMALS EQUALS 9
NO. OF CONTAMINATED EQUALS 1

	COL. B	COL. C	COL. D
	(X 10E8)	(X 10E0)	(X 10E-8)
MEAN	12.83	88.67	7.21
RANGE	12.95	78.00	4.61
MAX	21.42	142.00	9.20
MIN	8.47	64.00	4.58

. NO OUTLIERS

STOP
SRU'S:.6
!SWOTCH IN\$:MC656
!SAL

HOST MEDIATED ASSAY REPORT SHEET (REPEAT)

COMPOUND: FDA 71-36 ORGANISM: SALMONELLA G-46

DOSE LEVEL: HIGH - 5000 MG/KG

TREATMENT: IN VIVO, ORAL, SUBACUTE DATE STARTED: JUNE 13, 1973

ANI MAL NUMBER	A RAW CFU X 10E7/0.6ML.	B TOTAL CFU X 10E8/1.0ML	C TOTAL NO. MUTANTS X 10E0/1.0ML	D MUTATION FRE (C/B) X 10E-8
1	43.90	7.32	44.00	6.01
2	47.40	7.90	41.00	5.19
3	53.60	8.93	42.00	4.70
4	70.10	11.68	28.00	2.40
5	39.30	6.55	15.00	2.29
6	51.50	8.58	36.00	4.19
7	42.80	7.13	41.00	5.75
8	61.50	10.25	64.00	6.24

NO. OF ANIMALS EQUALS 8
NO. OF CONTAMINATED EQUALS 2

	COL. B	COL. C	COL. D
	(X 10E8)	(X 10E0)	(X 10E-8)
MEAN	8.54	38.88	4.60
RANGE	5.13	49.00	3.95
MAX	11.68	64.00	6.24
MIN	6.55	15.00	2.29

NO OUTLIERS

STOP SRU'S:.6 !SWITCH IN\$:MC654 !SAL !SWITCH IN\$:MC654 !SAL !OFF

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TERMATE PRINT FILE
  _RUn_ 100+(FU007, ______60,10006 -
G NUMBER= 200 .
P PAP+5 PRINT OR SIX-PART PAPER
        HOST MEDIATED ASSAY REPORT SHEET
       COMPOUND: FDA 71-36
                          ORGANISM: SACCHAROMYCES D+3
       DOSE LEVEL: HEGATIVE CONTROL - SALINE
       THEATMENT: IN VIVO, ORAL, ACUTE DATE STARTED: MAY 1, 1972
                            TOTAL CHU
                                        TOTAL
                                                 RECOMBICEU
       RECOMBINANTS SCREENED X
       NUMBER
                10E5/1.0ML
                           1005/1.0所
                                       /1.0例上
                                                   105-5
           422.00
                                        .....3.00
                                                      7.11
                  173+00
                                .17
                                           1.00
                                                      5.78
                  133.00
                                زله
                                           • 90
                                                      • 00
                648.40 .65
                                         4.00
                                                    6.17
                              .27
                  266+90
                                         2.00
                                                      7.52
        6
                  109.60
                               .11
                                          1.00
                                                      9.17
        7 198.00 .20
                                          .00
                                                      • 99
                  371.00
                                .37
                                           1.00
                                                      2.70
                  181.00
                                ·18
                                                       • 90
       TOTAL ?
                              2.50
                                         12.00
       INO. OF ANIMALS EQUALS 9 9
       NO. OF CONTAMINATED EQUALS 1
T
                         4.80
       MEAN C/MEAN B =
                            COL . B
                                        ... COL. C
                                                   COL. D
                              (X 10:5)
                                         (x 10E0)
                                                    (X 108-5)
                   MEAN
                                 .28
                                          1.33
                                                    4.37
                   RANCE _
                                           4.0
                                 ...54
                                                      9.17
                   MAX
                                           4.00
                                 • 05
                                                      9.17
                   MIN
                                 .11
                                            •06
                                                      . .05
       NO OUTLILES
```

COM	POUNC: FD	a 71-36		ORGANISM: SAG	CCHAROMYCES E
998	E LEVEL:	POSITIVE CO	DETROL - EMS 35		e e e e e e e e e e e e e e e e e e e
THE	ATMENT: I	VIVO. OR:	A ACUTE	DATE STARTED	e sand e engan
,		The state of the s	1 W. F	PRIL SIPRILL	 BBN LABOUR MAG
		ħ.	Ð		
	ti. Min titi kalen oran ayayya yancasa carr		TOTAL CFU	TOTAL	RECOMB/CFU
ANI	and the second s	RAW CEU X	SCREENED X	RECOMBINANTS	SCHEENED X
NUM	CER 1	0 <u>6</u> 5/1.0ML	10E5/1.0ML	/1.UHL	100-5
1		414.90	•41	2 t 0 0	FO 73
Ž.		283.00		21.00 18.00	50 • 73
3		301.00	•30	20.60	65•61 66•45
4		299.60	• 30	14.40	
5		232.00	• 23	11.00	46.82 47.41
6		326.00	•33	19.00	58+28
7		103.90	•10	ತ್ತು00	77.67
&		260+00			77•07 56•39
9		193.00	•19	12.00	62.18
10		392.00	•39	22.00	56.12
			. را سوی و داران درسانید ومد انسان دارد. سوی و داران درسانید ومد انسان دارد.		e egenement a la companya de la comp
TOTA	/_		2.61	160.00	
•		S EQUALS		100.00	The second secon
NO•.	OF ANIMAL	S EQUALS	19	150.00	
NO•	OF ANIMAL		10 6.95 COL. B	CoL. E	cot. D
NO•	OF ANIMAL	3. =	10 6.96 COL. B (X 10,5)	COL. ((% 19E0)	(X 107-5)
NO•	OF ANIMAL	MEAN	COL. B (X 1025)	COL. C (% 19E0) 16.00	(X 107-5) 58.57
NO•	OF ANIMAL	MEAN RANGE	COL. B (X 10,5) .28 .31	COL. C (X 10E0) 16.00 14.00	(X 107-5) 58.57 30.85
NO•	OF ANIMAL	MEAN RANGE MAX	COL. B (X 10,5) .29 .31	COL. C (X 10E0) 16.00 14.00 22.00	(X 10T-5) 58.57 30.85 77.67
NO•	OF ANIMAL	MEAN RANGE	COL. B (X 10,5) .28 .31	COL. C (X 10E0) 16.00 14.00	(X 107-5) 58.57 30.85
NO•	OF ANIMAL	MEAN RANGE MAX MIN	COL. B (X 10,5) .25 .31 .41	COL. C (X 10E0) 16.00 14.00 22.00	(X 10T-5) 58.57 30.85 77.67 46.82
NO•.	OF ANIMAL	MEAN RANGE MAX MIN	COL. B (X 10,5) .25 .31 .41	COL. C (X 10E0) 16.00 14.00 22.00 8.00	(X 10T-5) 58.57 30.85 77.67 46.82
NO.	OF ANIMAL	MEAN RANGE MAX MIN	COL. B (X 10,5) .25 .31 .41	COL. C (X 10E0) 16.00 14.00 22.00 8.00	(X 10T-5) 58.57 30.85 77.67 46.82
NO.	OF ANIMAI	MEAN RANGE MAX MIN	COL. B (X 10.5) •29 •31 •41 •10 SUMMARY WITH	COL. C (A 19E0) 16.00 14.00 22.00 8.00 OUTLIERS REMOVE	(X 10T-5) 58.57 30.85 77.67 46.82
NO.	OF ANIMAI	MEAN RANGE MAX MIN	COL. B (X 10,5) .25 .31 .41 .10 SUMMARY WITH	COL. C (% 1980) 16.00 14.00 22.00 8.00 OUTLIERS REMOVE	(X 10T-5) 58.57 30.85 77.67 46.82
NO.	OF ANIMAI	MEAN RANGE MAX MIN	COL. B (X 10L5) .25 .31 .41 .10 SUMMARY WITH	COL. C (% 10E0) 16.00 14.00 22.00 8.00 OUTLIERS REMOVE (% 10E0) 16.69	COL. D (X 101-5) 58.57 30.85 77.67 46.82
NO.	OF ANIMAI	MEAN RANGE MAX MIN	COL. B (X 1025) .25 .31 .41 .10 SUMMARY WITH COL. B (X 1025) .22	CoL. C (% 19E0) 16.00 14.00 22.00 8.00 OUTLIERS REMOVE (% 10E0) 16.89 11.00	COL. D (X 107-5) 58.57 30.85 77.67 46.82 ID (X 107-5) 56.84 19.62
ME AI	OF ANIMAI	MEAN RANGE MAX MIN	COL. B (X 10L5) .25 .31 .41 .10 SUMMARY WITH	COL. C (% 10E0) 16.00 14.00 22.00 8.00 OUTLIERS REMOVE (% 10E0) 16.69	COL. D (X 101-5) 58.57 30.85 77.67 46.82

TREATMENT: IN VIVO. ORAL. ACUTE DATE STARTED: MAY 1. 1972

.34

.11

.74

•22

TOTAL CFU

SCREENED X

NUMBER 10E5/1.0ML 10E5/1.0XL /1.0ML

ORGANISM: SACCHAROMYCES D-3

RECOMB/CFU

SCREENED X

102-5

14.66

28.04

6.83

6.73

9.17

TOTAL

RECOMBINANTS

3.00

2.00

5.00

... 2.00

COMPOUND: FDA 71-35

AHIMAL

DOSE LEVEL: LOW - 50 MG/KG

RAH CHU X

341.00

293.00

743.00

218.00

107.00

	7	250 • ii	^	♦ 1-	5•00	7.75
	8	178•0 603•0		•15 •69	1.00	5.62 9.09
	ò	477.0		•4b	6.00 4.00	8.39
	_TOTAL .	en anne de la companie de la compani	van en managan en en en en	3.22		·
	NO. OF	ANIMALS EQUA	LS 9		•	
		DEAD ANIMALS		1	and the section of th	
				*	•	
	MEAN CA	MEAN B =	9.32			
		•		col. B	CCL. C	COL. D
	· ·	man or some and an orall or some and an orall or some	e age of the contract of the c	(X 10L5)	(X 10E0)	13 (X 105-5)
		MEAN		• 36,	3.33	19,79
		RANGE	•	• 64	5.00	55.45
			-			
no district de la companya de district		XAM			6.00	28.04
			The second secon		6.00 1.00	28.04 5.62
		XAM		•74	1.00	5,62
		XAM		•74		5.62
		XAM		•74	1.00	5,62
	MEAN CA	XAM		•74	1.00	5,62
	MEAN C	NIM	* SUMA	•74	1.00	5.62 D
	MEAN C	MEAN B =	* SUMA	•74 •11 MARY WITH O	1.00 UTLIERS REMOVE Col. C (x 10E0)	5.62 CoL. 0 (x 107-4
	MEAN, C	MEAN B =	* SUMA 8.68	•74 •11 MARY WITH OF COL• 8 (X 10:5) •39	COL. C (X 10E0) 3.37	5.62 D CoL. D (x 107-4 8.66
	MEAN CA	MEAN B = MEAN KANGE	* SUMA 8.68	•74 •11 MARY WITH OF COL• 8 (X 10:5) •39 •36	COL. C (X 10E0) 3.37 5.00	COL. D (X 107-4 8.66 9.64
	MEAN CA	MEAN B =	* SUMA 8.68	•74 •11 MARY WITH OF COL• 8 (X 10:5) •39	COL. C (X 10E0) 3.37	5,62

	cc»Pound:	FDA 71-36		ORGANISM: SA	CCHAROMYCES D-
f !	DOSE LEVEL	: INTERMEDIAT	E - 500 MG/KG		
A	TREATMENT:	IN VIVO. ORA	L. ACUTE	DATE STARTED	MAY 1, 1972
r		<u> </u>		C	D
r	ANIMAL NUMBER	RAW CFU X 10E5/1.0ML	TOTAL CFU SCREENED X 1065/1.05L	TOTAL RECOMBINANTS /1.OML	RECOMBICEU SCHEENED X 10[-5
<u>^</u>	.1	103.00 233.00	.16	1.00	9•71 8•53
f ·	3 4 5	145.00 742.00 433.00	•14 •74 •43	2.00 4.00	13.79 5.39
6	6 7 8	367.00 288.00	•37 •29	3.00 3.00 3.00	2•31 8•17 10•42
	9	243.00 421.00	•24 •42	1.00 2.00 5.00	7.61 8.23 11.83
	TOTAL.	The second section of the second section of the second section of the second section s	3.11	₹ 4•₽0	
	Nu. of Ani	ALS EQUALS	10	and organization of the second organization organi	And the second s
	MEAN C/HEAN	1 B =	7.72		· · · · · · · · · · · · · · · · · · ·
			COL. 9 (X 1025)	COL. C (X 10E0)	COL. D (X 10E-5)
		MEAN RANGE MAX	•31 •64 •74	2.40 4.00	8.59 11.48
	NO OUTLIERS	MIN	•10	1.00	13.79 2.31

		FDA 71-36		OKONNIZW. 240	CHAROMYCES (
	DOSE LEVE	L: H1GH - 5000	MG/KG		the second secon
	TREATMENT	: IN VIVO. ORA	L. ACUTE	DATE STARTED	. MAY 1. 1972
neral mercer in		A		·	
			TOTAL CFU		RECOMP/CFI
		RAN CFU X	· · · · · · · · · · · · · · · · · · ·	RECOMBINANTS	
	NUMBER	10F5/1.0ML	10E5/1.0ML	/1.UML	102-5
	1	210.00	.21	8.00	38.19
	2			5.00	
	3	289.00	424	5.00	17.30
	ų	193.00	•19	7.00	36.27
		302.00	30		9.93
	Ď	280.00	•25	6.00	21.43
	7	169.00	.17	2.00	11.93
		424.00	42		
	9	163.00	•16	1.00	6.13
	TOTAL		2.38	41.00.	.4
		NIMALS EQUALS ONTAMENATED FRO			
months and a second		VIMALS EQUALS PATAMINATED EQU			e
	NO. OF CO		ALS 1		
	NO. OF CO	UDB DEFAMIRATHO	7.82		
	NO. OF CO	UDB DEFAMIRATHO	7.82 COL. B	Col. C	
	NO. OF CO	UDB DBFANIKATHO	7.82 COL. B (X 10.5)	(X 10E0)	COL. D (X 105-5
	NO. OF CO	UDB DETAMIRATHO	7.82 COL. B (X 101.5)	(x 10E0) 4.56	(X 10E-5 18.76
	NO. OF CO	MEAN BEAN RANGE	7.82 COL. B (X 1015) .26 .26	(X 10E0) 4.56 7.00	(X 10E-5 18.76 31.06
	NO. OF CO	MEAN BEAN RANGE MAX	7.82 COL. B (X 101.5) .26 .26 .46 .92	(X 10E0) 4.56 7.00 8.00	(X 10E-5 18.76 31.06 39.10
	NO. OF CO	MEAN B = 1 MEAN RANGE MAX MIN	7.82 COL. B (X 1015) .26 .26	(X 10E0) 4.56 7.00	(X 10E-5 18.76 31.06
	MEAN CYME	MEAN B = 1 MEAN RANGE MAX MIN	7.82 COL. B (X 101.5) .26 .26 .46 .92	(X 10E0) 4.56 7.00 8.00	(X 10E-5 18.76 31.06 38.10

TREATMENT: IN VIVO, ORAL, ACUTE DATE STARTED: MAY 5, 1972

.24

TOTAL CFU

SCREENED X

DOSE LEVEL: NEGATIVE CONTROL - SUBACUTE TRIALS

RAW CFU X

243.00

428.00

193.00

207.00

ORGANISM: SACCHAROMYCES D-3

SCREENED X

RECOMBINANTS

1.00

2.00

1.00

10E5/1.0ML /1.0ML 10E-5

COMPOUND: FDA 71-36

NUMBER 10E5/1.9ML

ANIMAL

6 333.00 .33 1.00	3.60
6 333.00 .33 1.00 7 186.00 .19 .00	• 78
8 291.00 .29 2.00	6.37
9 411.00 .41 3.00	7.30
10 382.00 .38 1.00	2.62
TOTAL 3.02 14.00	er velte er research en
NO. OF ANIMALS EQUALS 10	
MEAN C/MEAN B = 4.63	
COL. 8 COL. C (X 1055) (X 1080)	col. n (x 105-5)
MEAN .30 1.40	4.43
RANGE •24 3.09	7.30
MAX •43 3•00	7.30
MIN •19 •00	•00
* SUMMARY WITH OUTLIERS REMOV	VED
MEAN C/MEAN B = 4.93	
COL. IS COL. C	col. n
$(\chi 10; S) \qquad (\chi 10E0)$	(X 10E-5)
1.56	4.02
RANGE •23 2.00	4.68
FAX •43 3.03	7.00

COMPOUND	: FDA 71-35	'	ORGANISM: SAC	CHAROMYCES D
DOSE LEV	EL: LOW - 50	MG/KG		
TREATMEN	T: IN VIVO, ORA	L. SUBACUTE	DATE STARTED	MAY 5, 1972
	Δ	3		
AHIMAL NUMBER	RAW CFU X 10E5/1.OAL	TOTAL CFU SCREENED X 10E5/1.0ML	TOTAL RECOMBINANTS	RECOMB/CFU SCREENED X
	TOPOS X COME	TACE OF TACORE	/1.0ML	105-5
1	208.00	.21	1.00	4.81
	167.00	• 17		06. _•
3	370.00	•37	1.00	2.70
` 4	771.00	• 77	3.00	3.49
	728.00	73	7.00	<u>9•62</u>
- 6 7	361.00	•36	2.00	5.54
	518.00	•52	2.00	3∙ 8€
9	222.00			4.52
.10	438.00 720.00	•44 •76	2.00 3.00	4.57 4.17
TOTAL	- 	enversionemen et enversionemen et en la variable et en enversionemen et en enversionemen et en enversionemen e	en semanen servan er i sis sis segiste	and a a o o o o o o o o o o o o o o o o o o
	NIMALS FORMS	4.50 10	22.00	
	NIMALS EQUALS			••• <u>•</u> ••••••••••••••••••••••••••••••••
	•			
	•	10 		
	•	10 4.69 COL. B	CoL. C	cot. D
	EAN B =	COL. 3 (X 1015)	CoL. C (X 10E0)	(X 10E-5)
	EAN B =	COL. B (X 1015)	CoL. C (X 10E0) 2.20	(X 105-5) 4.37
	EAN B =	COL. B (X 1015) .45	CoL. C (X 10E0) 2.20 7.80	1 (X 10E-5) 4.37 9.62
	EAN B =	COL. B (X 1015)	CoL. C (X 10E0) 2.20	(X 105-5) 4.37
	EAN B = HEAN RANGE HAX	COL. B (X 10L5) .45 .00	CoL. C (X 10E0) 2.20 7.00 7.00	(X 105~9) 4.37 9.62 9.62
	EAN B C MEAN RANGE MAX MIN	COL. B (X 10L5) •45 •00 •77 •17	CoL. C (X 10E0) 2.20 7.00 7.00	(X 105-9) 4.37 9.62 9.62
	EAN B C MEAN RANGE MAX MIN	COL. B (X 10L5) •45 •00 •77 •17	CoL. C (X 10E0) 2.20 7.00 7.00	(X 105~9) 4.37 9.62 9.62 .00
NO. OF A	EAN B S MEAN RANGE MAX MIN	COL. B (X 10L5) •45 •00 •77 •17	CoL. C (X 10E0) 2.20 7.00 7.00	(X 105-5) 4.37 9.62 9.62
NO. OF A	EAN B C MEAN RANGE MAX MIN	COL. B (X 10L5) .45 .50 .77 .17 SUMMARY WITH	COL. C (X 10E0) 2.20 7.00 7.00 .60 COL. C	(X 105-9) 4.37 9.62 9.62 .00
NO. OF A	EAN B =	COL. B (X 10L5) .45 .00 .77 .17 SUMMARY WITH 4.16 COL. B (X 10.5)	COL. C (X 10E0) 2.20 7.00 7.00 .00 OUTLIERS REMOVE	(X 10F-5) 4.37 9.62 9.62 .00
NO. OF A	EAN B = AEAN RANGE MAX MIN	COL. B (X 10L5) .45 .50 .77 .17 SUMMARY WITH	COL. C (X 10E0) 2.20 7.00 7.00 .60 OUTLIERS REMOVE	COL. D (x 105-5)
NO. OF A	MEAN RANGE MAX MIN	COL. B (X 10L5) •45 •00 •77 •17 SUMMARY WITH 4.16 COL. B (X 10L5) •45 •6	CoL. C (X 10E0) 2.20 7.00 7.00 .00 OUTLIERS REMOVE (X 10E0) 1.88 2.00	COL. D (x 105-5) 9.62 9.62 .00 (x 105-5) 9.25 2.64
NO. OF A	EAN B = AEAN RANGE MAX MIN	COL. B (X 10L5) .45 .00 .77 .17 SUMMARY WITH 4.16 COL. B (X 10.5)	COL. C (X 10E0) 2.20 7.00 7.00 .60 OUTLIERS REMOVE	COL. D (x 105-5 9.62 9.62 .00 (x 105-5 4.25

	CORPOUND	FDA 71-36		ORGANISM: SAC	CHAROMYCES D-3
	DOSE LEVE	L: INTERMEDIA	TE - 500 MG/KB		* • • • • • • • • • • • • • • • • • • •
	TREATMEN	F: IN VIVO, OR	AL. SUBACUTE	DATE STARTED	MAY 5, 1972
			B TOTAL CFU	C	D COURT COURT
	ANIMAL NUMBER	RAW CFU X 10EU/1.0ML	SCREENED X 1		
	1	262.00	•23	2.00	7.09
A		198•00 450•00 227•00		1.00 1.00 1.00	5.05 2.23 4.41
.,	5 6	284 • 00 457 • 00	• 28 • 45	1.00 3.00	3.52 6.56
	7 	- 493•00 256•00	•49 •26	2.60 1.00	4.06 3.91
	9 19	283•00 286•00	•28 •29	1.00 1.00	3. 53 3. 50
	TOTAL		3.22	14.90	 A section is related to the control of the control of
<u> </u>	NO. OF A	HMALS EQUALS	10		en e energia de la company de
	MEAN C/A	EAN B =	. 4.35	enter de la companya del companya del companya de la companya de	
	mine a similar and make a similar second and similar second and second and second and second and second and second		COL. B (X 1025)	COL. C (X 10E0)	COL. D (X 105-5)
		MEAN RANGE	•32 •30	1.40 2.00	4.38 4.87
	NO OUTLI	MAX MIN LRS	•49 •20	3.00 1.00	2,22

	COMPOUND:	FDA 71-36	ORGANISM: SACCHAROMYCES DA		
	DOSE LEVE	L: HIGH - 5000	MG/KÇ	n en	
	TREATMENT	: IN VIVO. ORA	L. SUBACUTE	DATE STARTED	1.8AY 51 1972
		A		<u> </u>	D
	ANIMAL NUMBER	RAN CFU X 10E3/1.GML	TOTAL CFU SCREENED X 1065/1.0%L	TOTAL RECOMBINANTS /1.UML	RECOMBICER SCREENED X 105-5
	. 1 2	352.60 280.00	•35 •22 ::	1.00	2.85 3.57
	3 4 5	297.60 153.00 188.00	•36 •16 •19	1.00 .00 2.00	3.37 -90 10.64
	5 7 8	285+90 483+90 326+90	•29 •46 •35	2.00 2.00 4.00	6.99 8.28 3.07
1	ŋ	239.00	•24	2.00	8.37
	Tural		2.61	14.00	
· · · · · · · · · · · · · · · · · · ·		HIMALS EQUALS REERED OUT OF R	ANGE, EQUALS	1	
· · · · · · · · · · · · · · · · · · ·	MEAN C/ME	an s =	5.37	en e	
	energy on the company of the company		COL. B (X 1015)	CoL. C (x 10E0)	CCL. D (X 10E~5)
		MEAH RANGE MAX	•29 •32 •46	1.50 4.00 4.00	5.24 10.64 10.64
	NO OUTLIE	MIN	•16	•00	.00

3. Cytogenetics

a. <u>In vivo</u>

(1) Acute study

The chromosomal abnormalities observed in the positive controls were significantly higher than either the negative controls or the compound. An increased frequency of breaks was observed in the high dosage level, but these were not significantly different from the negative control values. The negative control values are within normal limits. Mitotic indices in the negative control and the compound test groups were within normal values.

(2) Subacute study

Mitotic indices and breaks were within.

normal values for all groups in this study.

b. <u>In vitro</u>

Anaphase preparations were examined in this test.

The positive control compound produced a significantly higher percentage of aberrations on the chromosomes than the negative control or the test compound. Depression of the mitotic index due to the positive control was not as pronounced as in the <u>in vivo</u> test. The compound did not produce any observed aberrations. The negative control values were normal.

c. CYTOGENETIC SUMMARY TABLES

CONTRACT FDA 71-268

COMPOUND FDA 71-36

CALCIUM PROPIONATE



COMPOUND FDA 71-36 CALCIUM PROPIONATE ACUTE STUDY METAPHASE SUMMARY SHEET

Compound	Dosage (mg/kg)	Time*	No. of Animals	No. of Cells	Mitotic*** Index %	% Cells with Breaks	% Cells with Reunion	% Cells Other Aber.**	% Cells with aber.
Negative Control	saline	6 24 48	3 3 3	150 150 150	8 9 12	2 0 2	0 0 0	0 0 0	2 0 2
Low Level	50	6 24 48	5 5 5	250 250 250	13 13 12	1 0 0	0 0 0	0 0 0	1 0 0
Intermediate Level	500	6 24 48	5 5 5	250 250 250	10 9 6	0 0 3	0 0 0	0 0 0	0 0 3
High Level	5000	6 24 48	5 5 5	250 250 250	12 8 11	4 3 2	0 0 0	0 0 0	4 3 2
Positive Control TEM	0.3	48	5	250	4	33	12	6 a	48

^{*} Time of kill after injection (hours).

** Cells that have polyploidy (P), pulverization (pp), or greater than 10 aberrations (a).

***% of cells in mitosis: 500 cells observed/animal.

COMPOUND FDA 71-36 CALCIUM PROPIONATE SUBACUTE STUDY METAPHASE SUMMARY SHEET

Compound	Dosage* (mg/kg)	No. of Animals	No. of Cells	Mitotic*** Index %	% Cells with Breaks	% Cells with Reunion	% Cells Other Aber.**	% Cells with aber.
Negative Control	SALINE	3	150	10	0	0	0	. 0
Low Level	50	5	250	10	1	0	0	1
Intermediate Level	500	5	250	9	. 0	0	0	0
High Level	5000	5	250	12	0	0	0	0

^{*} Dosage 1X/day X 5 days.
** Cells that have polyploidy (P), pulverization (pp), or greater than 10 aberrations (a).
*** % of cells in mitosis: 500 cells observed/animal.

COMPOUND FDA 71-36 CALCIUM PROPIONATE ANAPHASE SUMMARY SHEET

Compound	Dosage (mcg/ml)	Mitotic ** Index	No. of Cells	% Cells with Acentric Frag.	% Cells with Bridges	% Multipolar Cells	% Cells Other Aber.*	% Cells with aber.
Low Level	0.4	6	100	0	0	0	0	0
Medium Level	4	3	100	0	0	0	0	. 0
High Level	40	2	100	0	0	0	0	0
Negative Control	saline	4	100	0	0	0	0	0
Positive Control (TEM)	0.1	2	100	6	2	0	0	8

^{*}Cells that have polyploidy (P), pulverization (pp), or greater than 10 aberrations (a). **% of cells in mitosis: 200 cells observed/dosage group.

4. Dominant Lethal Assay

The interpretation of these data were made by Dr. David Brusick, Assistant Professor of Microbiology, Howard University, Washington, D.C., as a consultant to Litton Bionetics, Inc.

Fertility Index

- Acute Negative control all treated groups in week 7 showed significant decreases.
- Subacute Negative control significant differences were found at all three doses on week 7. Other significant differences were found at some doses at weeks 2 and 6.

Average Number of Implantations per Pregnant Female

- Acute Negative control significant dose-related decreases were observed in weeks 3 and 4 at the high dose level.
- Subacute Negative control significant, dose-related, decreases were observed in week 2.

Average Number of Corpora Lutea

- Acute Negative control significant decreases were observed in weeks 3, 6 and 8. All three doses showed significant decreases during week 8. Dose-relationship noted in weeks 6 and 8.
- Subacute Negative control significant, dose-related, decrease in week 2.

Average Preimplantation Losses per Pregnant Female

- Acute Negative control significant, dose-related, increases in weeks 3 and 7. Significant, dose-related, decreases in week 8.
- Subacute Negative control significant, dose-related, increases in weeks 6 and 7.



Average Resorptions per Pregnant Female

Acute - Negative control - significant, dose-related, increases at weeks 1, 2 and 6.

Subacute - Negative control - no significant findings.

Proportion of Females with One or More Dead Implantations

Acute - Negative control - no significant dose-related effects.

Subacute - Negative control - no significant increases of females with one or more dead implants. Some weeks (1 and 4) showed significant decreases.

Proportion of Females with Two or More Dead Implantations

Acute - Negative control - no significant effects.

Subacute - Negative control - no significant effects.

Dead Implants/Total Implants

Acute - Negative control - significant increase at high dose in week 1.

Subacute - Negative control - no significant effects.



DOMINANT LETHAL SUMMARY TABLES

CONTRACT FDA 71-268

COMPOUND FDA 71-36

CALCIUM PROPIONATE



TABLE I

COMPOUND 36

STUDY ACUTE

FERTILITY INDEX

ARITH DOSE	WEEK	HISTORICAL CONTROL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL . 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG	POSITIVE CONTROL
•	1,	109/159=0.69	13/20=0.65	14/20=0.70	17/19=0.90	13/20=0.65	10/18=0.56
•	2	119/159=0.75	16/19=0.85	17/20=0.85	18/20=0.90	15/20=0.75	11/19=0.58
	, 3	119/158=0.76	16/20=0.80	19/20=0.95	19/20=0.95	19/20=0.95	11/20=0.55
	4	136/160=0.85	18/20=0.90	18/20=0.90	19/20=0.95	16/20=0.80	16/20=0.80
	5	127/159=0.80	19/20=0.95	20/20=1.00	18/19=0.95	17/20=0.85	18/20=0.90
•	б	128/159=0.81	13/19=0.69	19/20=0.95*	17/19=0.90	15/18=0.84	19/20=0.95*
•	7	133/157=0.85	20/20=1.00	14/20=0.70**	16/20=0.80*	12/18=0.67**	20/20=1.00
	8 .	133/160=0.84	18/20=0.90	15/19=0.79	15/19=0.79	14/18=0.78	16/20=0.80

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

ONE !,* = SIGNIFICANT AT P LESS THAN 0.05
TWO !.* = SIGNIFICANT AT P. LESS THAN 0.01

^{*} SIGNIFICANTLY DIFFERENT FROM CONTROL

[!] SIGNIFICANT LINEAR RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

COMPOUND 36

STUDY ACUTE

AVERAGE NUMBER OF IMPLANTATIONS PER PREGNANT FEMALE

TABLE II

G S E	ARIT		WEEK	HISTORICAL CONTROL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG	POSITIVE CONTROL
			1	1351/109=12.4	171/13=13.2	162/14=11.6	199/17=11.7	168/13=12.9	52/10= 5.2**ลอเ **ลอเ
1			2	1427/119=12.0	183/16=11.4	175/17=10.3 @D	232/18=12.9 @I	191/15=12.7 I	86/11= 7.8*an **aai
1 1	٤!		3	1435/119=12.1		232/19=12.20D aai	239/19=12.6	221/19=11.6**@@I	D 93/11= 8.5**ออเ *อบ
	8 1	\$.	4	1626/136=12.0	222/18=12.3	232/18=12.9	229/19=12.1	177/16=11.1aD	171/16=10.7**aat **aat
			5	1466/127=11.5	221/19=11.6	231/20=11.6	201/18=11.2	212/17=12.5 aī	205/18=11.4
		•	6	1512/128=11.8	138/13=10.6	214/19=11.3	188/17=11.1	178/15=11.9	167/19= 8.8aD **aar
		•	7	1626/133=12.2	252/20=12.6	162/14=11.6	185/16=11.6	134/12=11.2	205/20=10.3*@aD **@ar
	,		8	1551/133=11.7	213/18=11.8	164/15=10.9	180/15=12.0	156/14=11.1	205/16=12.8 *aI

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

E AND * = TWO-TAILED TEST ! AND @ = ONE-TAILED TEST

ONE !,&, $\hat{\omega}$,* = SIGNIFICANT AT P LESS THAN 0.05 TWO !,&, ω ,* = SIGNIFICANT AT P LESS THAN 0.01

^{*, 0} SIGNIPICANTLY DIFFERENT FROM CONTROL

^{8,!} SIGNIFICANT RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE III

COMPOUND 36

STUDY ACUTE

AVERAGE CORPORA LUTEA PER PREGNANT FEMALE

G SE	ARITH DOSE	WEEK	HISTORICAL CONTROL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG	POSITIVE CONTROL
	• .	1	1504/109=13.8	173/13=13.3	181/14=12.9	227/17=13.4	181/13=13.9	91/10= 9.1**@a **@a
!	•	2	1588/119=13.3	202/16=12.6	186/17=10.9an **ā	232/18=12.9	191/15=12.7	129/11=11.7 *@aD
!		. 3	1565/119=13.2	217/16=13.6	232/19=12.20D	239/19=12.6	238/19=12.5	116/11=10.6*@D *@D
	8 ! 1133	. 4	1784/136=13.1	225/18=12.5	232/18=12.9	234/19=12.3	180/16=11.3*ap **a	180/16=11.3*@@D **@@i
		5	1648/127=13.0	230/19=12.1 aD	241/20=12.1 aD	217/18=12.1	213/17=12.5	209/18=11.6 *aap
! !!	ε !!	6	1689/128=13.2	180/13=13.9	295/19=15.5 **a		0I 231/15=15.4 *@@I **@	213/19=11.2*aD aI *aaD
		7	1767/133=13.3	255/20=12.8	186/14=13.3	227/16=14.2	169/12=14.1	250/20=12.5
!!		8	1823/133=13.7	319/18=17.7 **@@		aD 199/15=13.3**	*@@D210/14=15.0@D @I	243/16=15.2

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

& AND * = TWO-TAILED TEST
! AND @ = ONE-TAILED TEST

ONE 1,8, ω ,* = SIGNIFICANT AT P LESS THAN 0.05 TWO 1,8, ω ,* = SIGNIFICANT AT P LESS THAN 0.01

^{*, @} SIGNIFICANTLY DIFFERENT FROM CONTROL

^{8,!} SIGNIMICANT RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE IV

COMPOUND 36

STUDY ACUTE

AVERAGE PREIMPLANTATION LOSSES PER PREGNANT FEMALE

	ARITH DCSE	WEEK	HISTORICAL CONTROL	NEGATIVE CONTROL		DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG	POSITIVE CONTROL
		1	153/109= 1.4	2/13= 0.2 **@@D		DI 28/17= 1.7*a	I 13/13= 1.0	39/10= 3.9**aa *aar
!!!	1133	2	161/119= 1.4	19/16 = 1.2	11/17= 0.7	0/18= 0.0 D **a	0/15= 0.0 aad **aa	43/11= 3.9*@I *@I
11	1133	3	130/119= 1.1	0/16= 0.0 **@@D	0/19= 0.0	0/19= 0.0 *aaD ***	17/19= 0.9*aai	23/11= 2.1*@@I
11	ε !	4	158/136= 1.2	3/18= 0.2 **@@D	0/18= 0.0	5/19= 0.3 *@@D ***	3/16= 0.2 aad **aa	9/16= 0.6 D
1:	8 !!	5	182/127= 1.4	9/19= 0.5 **@@D	10/20= 0.5	16/18= 0.9 aad	1/17= 0.1 **@@]	4/18= 0.2 **aa:
!!	1133	6	177/128= 1.4	42/13= 3.2 *@I	81/19= 4.3	91/17= 5.4aI *aaI **a	53/15= 3.5 00I **da	46/19= 2.4 I
	ε ! ε !!	7	141/133= 1.1	3/20= 0.2 **@aD			aai 35/12= 2.9**aa ai	I 45/20= 2.3**aa **aa
!	ε !!	8	272/133= 2.1	106/18= 5.9 **aai			0aD 54/14= 3.9 *aaI	

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

 ε AND * = TWO-TAILED TEST ! AND @ = ONE-TAILED TEST

ONE $1, \varepsilon, \omega, * = SIGNIFICANT AT P LESS THAN 0.05$ TWO !, &, a, * = SIGNIFICANT AT P LESS THAN 0.01

*, D SIGNIFICANTLY DIFFERENT FROM CONTROL

8,! SIGNIFICANT RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE V

COMPOUND 36

STUDY ACUTE

AVERAGE RESORPTIONS (DEAD IMPLANTS) PER PREGNANT FEMALE

G SE	ARITH DOSE	WEEK	HISTORICAL CONTROL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG	POSITIVE CONTROL
! !!	1133	1	28/109=0.26	3/13=0.24	6/14=0.43	6/17=0.36	13/13=1.00@I *@I	42/10=4.20**aai **aai
! !!	1133	2	53/119=0.45	4/16=0.25	15/17=0.89	47/18=2.62*@I @I	41/15=2.74aI aI	36/11=3.28**aai **aar
	. '	3	61/119=0.52	10/16=0.63	21/19=1.11	4/19=0.22 @D	4/19=0.22 aD	14/11=1.28 *@I
ī		4	62/136=0.46	12/18=0.67	4/18=0.23 ad	2/19=0.11an **aar	5/16=0.32	73/16=4.57**aai **aai
1!	ε!!	5	74/127=0.59	6/19=0.32	6/20=0.30	6/18=0.34	1/17=0.06 **@@D	16/18=0.89
11	!	6	58/128=0.46	2/13=0.16 *@D	5/19=0.27	17/17=1.000I	12/15=0.80*@I	23/19=1.22**@@I **@@I
11	ε !	7	65/133=0.49	1/20=0.05 **aaD	0/14=0.0 **@@I	1/16=0.07 **@@I	0/12=0.0 **aap	1/20=0.05 **aaD
		8	71/133=0.54	5/18=0.28	3/15=0.20 *aD	5/15=0.34	5/14=0.36	13/16=0.82

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

ε AND * = TWO-TAILED TEST
! AND ∂ = ONE-TAILED TEST

ONE 1,8, α ,* = SIGNIFICANT AT P LESS THAN 0.05 TWO 1,8, α ,* = SIGNIFICANT AT P LESS THAN 0.01

*, D SIGNIFICANTLY DIFFERENT FROM CONTROL

8, ! SIGNIFICANT RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE VI

COMPOUND 36

STUDY ACUTE

PROPORTION OF FEMALES WITH ONE OR MORE DEAD IMPLANTATIONS

ARIT DOSE		VEEK	HISTORICAL CONTROL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL . 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG	POSITIVE CONTROL
 !		. 1	24/109=0.23	3/13=0.24	3/14=0.22	6/17=0.36	7/13=0.54	8/10=0.80** **
		2	38/119=0.32	3/16=0.19	7/17=0.42	8/18=0.45	7/15=0.47	7/11=0.64*
	•	3	39/119=0.33	5/16=0.32	5/19=0.27	3/19=0.16	3/19=0.16	7/11=0.64
		4	46/136=0.34	6/18=0.34	3/18=0.17	2/19=0.11	4/16=0.25	14/16=0.88**
!	•	5	45/127=0.36	4/19=0.22	5/20=0.25	4/18=0.23	1/17=0.06	8/18=0.45
		6	44/128=0.35	2/13=0.16	3/19=0.16	7/17=0.42	8/15=0.54*	13/19=0.69**
:		7	46/133=0.35	1/20=0.05	0/14=0.0	1/16=0.07	0/12=0.0	1/20=0.05
		8	50/133=0.38	4/18=0.23	3/15=0.20	4/15=0.27	5/14=0.36	6/16=0.38

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

CNE !, * = SIGNIFICANT AT P LESS THAN 0.05
TWO !, * = SIGNIFICANT AT P LESS THAN 0.01

^{*} SIGNIFICANTLY DIFFERENT FROM CONTROL

[!] SIGNIFICANT LINEAR RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE VII

COMPOUND 36

STUDY ACUTE

PORPORTION OF FEMALES WITH TWO OR MORE DEAD IMPLANTATIONS

)G)SE	ARITH DOSE	WEEK	HISTORICAL CONTROL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVFL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG	POSITIVE CONTROL
•	!!	1	3/109=0.03	0/13=0.0	1/14=0.08	0/17=0.0	3/13=0.24	7/10=0.70**
	•	2	14/119=0.12	1/16=0.07	4/17=0.24	5/18=0.28	4/15=0.27	7/11=0.64**
	•	. 3	17/119=0.15	4/16=0.25	3/19=0.16	1/19=0.06	1/19=0.06	5/11=0.46
		4	12/136=0.09	3/18=0.17	1/18=0.06	0/19=0.0	1/16=0.07	13/16=0.82**
		5	18/127=0.15	2/19=0.11	1/20=0.05	2/18=0.12	0/17=0.0	4/18=0.23
		6	13/128=0.11	0/13=0.0	2/19=0.11	4/17=0.24	3/15=0.20	6/19=0.32*
		7	14/133=0.11	0/20=0.0	0/14=0.0	0/16=0.0	0/12=0.0	0/20=0.0
		8	18/133=0.14	1/18=0.06	0/15=0.0	1/15=0.07	0/14=0.0	4/16=0.25

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

CNE ! * = SIGNIFICANT AT P LESS THAN 0.05 TWO ! * = SIGNIFICANT AT P LESS THAN 0.01

^{*} SIGNIFICANTLY DIFFERENT FROM CONTROL

[!] SIGNIFICAN' LINEAR RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE VIII COMPOUND 36

STUDY ACUTE

DEAD IMPLANTS / TOTAL IMPLANTS

WFEK	HISTORICAL CONTROL	NEGATIVE I	DOSE LEVEL 50.000 MG/KG		DOSE LEVEL 5000.000 MG/KG	POSITIVE CONTROL
1	28/1351=0.03	3/171=0.02	6/162=0.04	6/199=0.04	13/168=0.08*@I *@I	42/ 52=0.81**@ **@@
2	53/1427=0.04	4/183=0.03	15/175=0.09	47/232=0.21@I @I	41/191=0.22aI aI	36/ 86=0.42**@ **@@
3	61/1435=0.05	10/217=0.05	21/232=0.10	4/239=0.02 *@@	,	14/ 93=0.16*aI *aI
4	62/1626=0.04	12/222=0.06	4/232=0.02 *@D	2/229=0.01	5/177=0.03 aad	73/171=0.43**@ **@@
5	74/1466=0.06	6/221=0.03 *ap	6/231=0.03	6/201=0.03	1/212=0.01 **@aD	16/205=0.08@1
6	58/1512=0.04	2/138=0.02	5/214=0.03	17/188=0.10	12/178=0.07	23/167=0.14**@/ **@@
7	65/1626=0.04	1/252=0.01 **aan	0/162=0.0 D **@	1/185=0.01 PD **a	0/134=0.0	1/205=0.01ai **aa
8	71/1551=0.05	5/213=0.03 @D	3/164=0.02	5/180=0.03	5/156=0.04	13/205=0.07

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT DIFFERENCES USING THE HISTORICAL CONTROL GROUP

^{* =} TWO-TAILED TEST

a = ONE-TAILED TEST

ONE *.0 = SIGNIFICANT AT P LESS THAN 0.05 TWO *.0 = SIGNIFICANT AT P LESS THAN 0.01

^{*,} a SIGNIFICANTLY DIFFERENT FROM CONTROL

TABLE I

COMPOUND 36

STUDY SUBACUTE

FERTILITY INDEX

E	ARITH DOSE WF	ΞK	HISTORICAL CONTROL	NEGATIVE CONTROL		DOSE LEVEL 50.000 MG/KG	DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG
	•	1	104/159=0.66	13/19=0.69	•	12/20=0.60	9/19=0.48	12/20=0.60
		2	118/160=0.74	14/20=0.70		14/20=0.70	19/20=0.95*	14/20=0.70
	•	3	119/159=0.75	12/19=0.64		14/19=0.74	15/20=0.75	14/19=0.74
		4	120/154=0.78	16/19=0.85		12/20=0.60	16/20=0.80	17/20=0.85
		5	122/157=0.78	18/20=0.90		19/20=0.95	17/19=0.90	18/20=0.90
	(6	136/159=0.86	18/19=0.95		11/20=0.55**	18/20=0.90	13/18=0.73
		7	135/155=0.88	19/19=1.00		9/19=0.48**	12/20=0.60**	12/19=0.64**

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

CNE ! * = SIGNIFICANT AT P LESS THAN 0.05
TWO ! * = SIGNIFICANT AT P LESS THAN 0.01

^{*} SIGNIFICANTLY DIFFERENT FROM CONTROL

[!] SIGNIFICANT LINEAR RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE II

COMPOUND 36

STUDY SUBACUTE

AVERAGE NUMBER OF IMPLANTATIONS PER PREGNANT FEMALE

G SE	DOSE	WEEK	CONTROL	CONTROL	50.000 MG/KG	500.000 MG/KG 50	000.000 MG/KG
!	ε!	1	1231/104=11.8	167/13=12.9	150/12=12.5	108/ 9=12.0	163/12=13.6 *@@I
:		2	1474/118=12.5	190/14=13.6 @I	178/14=12.7	222/19=11.7*aD	164/14=11.7aD
!	ε!	3 -	1405/119=11.8	139/12=11.6	155/14=11.1	181/15=12.1	181/14=12.9aI - *aI
		4	1414/120=11.8	191/16=11.9	137/12=11.4	180/16=11.3	195/17=11.5
		5 .	1462/122=12.0	219/18=12.2	237/19=12.5	199/17=11.7	231/18=12.8 @I
1 1	ε!	6	1626/136=12.0	179/18= 9.9 **@@	108/11= 9.8 D aD	183/18=10.2 *aD	132/13=10.2
		7	1566/135=11.6	204/19=10.7 @D	100/ 9=11.1	137/12=11.4	139/12=11.6

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

& AND * = TWO-TAILED TEST
! AND @ = ONE-TAILED TEST

ONE !, ε , ω , * = SIGNIFICANT AT P LESS THAN 0.05 TWO !, ε , ω , * = SIGNIFICANT AT P LESS THAN 0.01

*, @ SIGNIFICANTLY DIFFERENT FRCM CONTROL E,! SIGNIFICANT RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE III

COMPOUND 36

STUDY SUBACUTE

AVERAGE CORPORA LUTEA PER PREGNANT FEMALE

	ARITH DOSE	WEEK	HISTORICAL CONTROL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL D 500.000 MG/KG 5	OSE LEVEL 000.000 MG/KG
	!	1	1385/104=13.3	171/13=13.2	156/12=13.0	117/ 9=13.0	175/12=14.6
!! !!	!	2	1599/118=13.6	194/14=13.9	178/14=12.7	227/19=12.0*aa **a	D 173/14=12.4*aD aD *aD
:	1	· 3	1535/119=12.9	139/12=11.6 *aar	168/14=12.0	183/15=12.2	183/14=13.101
		4	1499/120=12.5	198/16=12.4	144/12=12.0	192/16=12.0	207/17=12.2
		5	1554/122=12.7	219/18=12.2	237/19=12.5	211/17=12.4	231/18=12.8
!	1	6	1809/136=13.3	193/18=10.7 **@@	130/11=11.8 D	220/18=12.20I	171/13=13.201
!!	ε !	7	1711/135=12.7	215/19=11.3 *aD	113/ 9=12.6	157/12=13.101	167/12=13.9*aaI

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

8 AND * = TWO+TAILED TEST
! AND @ = ONE-TAILED TEST

ONF 1,8,0,* = SIGNIFICANT AT P LESS THAN 0.05 TWO 1,8,0,* = SIGNIFICANT AT P LESS THAN 0.01

E.! SIGNIFICANT RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

^{*,} D SIGNIFICANTLY DIFFERENT FROM CONTROL

TABLE IV

COMPOUND 36

STUDY SUBACUTE

AVERAGE PREIMPLANTATION LOSSES PER PREGNANT FEMALE

OG OSE		RITH OSE	WEEK	HISTORICAL CONTROL		IVE D ROL	50.000		SE LEVE	L DOS MG/KG 500	E LEVEL	
· !		•	1	154/104= 1.	5 4/13=	0.3 **@@D		0.5 **@@n		1.0	12/12=	1.0
8!!			2	125/118= 1.	1 4/14=	0.3 **@@D	0/14=	0.0 **aap	5/19=	0.3 **@@D	9/14=	0.6
ε!!	ε	!	3	130/119= 1.	1 0/12=	0.0 **adD	13/14=	0.9	2/15=	0.1 **@@D		0.1 **aaD
			4	85/120= 0.	7 7/16=	0.4	7/12=	0.6	12/16=	0.8	12/17=	0.7
811	3	11	5	92/122= 0.	8 0/18=	0.0 **aaD		0.0 **@@D		0.7	0/18=	0.0 **@@D
6!! 6!!			6.	183/136= 1.	4 14/18=	0.8	22/11=	2.001	37/18=	2.1*aaI aI	39/13=	3.0*@@I *@I
5!! 5!!		!	7	145/135= 1.	1 11/19=	0.6	13/ 9=	1.401	20/12=	1.701	28/12=	2.3*aai *ai

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

E AND * = TWO-TAILED TEST ! AND @ = ONE-TAILED TEST

ONE $!, \varepsilon, \omega, *$ = SIGNIFICANT AT P LESS THAN 0.05 TWO $!, \varepsilon, \omega, *$ = SIGNIFICANT AT P LESS THAN 0.01

*, a SIGUIPICANTLY DIFFERENT FROM CONTROL

E,! SIGNIFICANT RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE V

COMPOUND 36

STUDY SUBACUTE

AVERAGE RESORPTIONS (DEAD IMPLANTS) PER PREGNANT FEMALE

OG OS E	ARITH DOSE	WFEK	HISTORICAL CONTRCL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG
		1	40/104=0.39	8/13=0.62	5/12=0.42	3/ 9=0.34	9/12=0.75 @I
1		2	59/118=0.50	18/14=1.29	33/14=2.36	49/19=2.58	16/14=1.15
6!!	ε 1	3	69/119=0.58	3/12=0.25	4/14=0.29	0/15=0.0 **@@I	1/14=0.08 **aaD
	1133	4	66/120=0.55	7/16=0.44	3/12=0.25	3/16=0.19 *ap	0/17=0.0 **aab **aab
		5	78/122=0.64	7/18=0.39	7/19=0.37 ap	12/17=0.71	7/18=0.39
1		6	62/136=0.46	•	15/11=1.37 aap	0/18=0.0 **aan	4/13=0.31
i		7	70/135=0.52	5/19=0.27	0/ 9=0.0 ad **adi	4/12=0.34	1/12=0.09 **@@D

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

& AND * = TWO-TAILED TEST
! AND @ = ONE-TAILED TEST

ONE !, &, \(\partial\), * = SIGNIFICANT AT P LESS THAN 0.05
TWO !, \(\epsilon\), * = SIGNIFICANT AT P LESS THAN 0.01

*, @ SIGNIFICANTLY DIFFERENT FROM CONTROL E,! SIGNEFICANT RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE VI

COMPOUND 36

STUDY SUBACUTE

PROPORTION OF FEMALES WITH ONE OR MORE DEAD IMPLANTATIONS

OG OSE	ARITH DOSE	WEEK	HISTORICAL CONTROL	NEGATIVE CCNTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG
	! !	1	31/104=0.30	7/13=0.54	3/12=0.25	1/ 9=0.12*	7/12=0.59
		. 2	38/118=0.33	7/14=0.50	7/14=0.50	6/19=0.32	4/14=0.29
! !	•	3	42/119=0.36	2/12=0.17	3/14=0.22	0/15=0.0 **	1/14=0.08
! !	!!	4	42/120=0.35	6/16=0.38	3/12=0.25	3/16=0.19	0/17=0.0 **
		5	54/122=0.45	5/18=0.28	4/19=0.22	4/17=0.24	6/18=0.34
I I	¥ *	6	43/136=0.32	1/18=0.06	2/11=0.19	0/18=0.0	2/13=0.16
		7	42/135=0.32	3/19=0.16	0/ 9=0.0	3/12=0.25	1/12=0.09

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT PELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

ONE ! * = SIGNIFICANT AT P LESS THAN 0.05 TWO ! * = SIGNIFICANT AT P LESS THAN 0.01

^{*} SIGNIFICANTLY DIFFERENT FROM CONTROL

[!] SIGNIFICANT LINEAR RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE VII

COMPOUND 36

STUDY SUBACUTE

PORPORTION OF FEMALES WITH TWO OR MORE DEAD IMPLANTATIONS

	ARITH DOSE	WEEK	HISTORICAL CONTROL	NEGATIVE CCNTRCL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG
		1	8/104=0.08	1/13=0.08	2/12=0.17	1/ 9=0.12	2/12=0.17
		2	10/118=0.09	4/14=0.29	3/14=0.22	4/19=0.22	3/14=0.22
	•	. 3	17/119=0.15	1/12=0.09	1/14=0.08	0/15=0.0	0/14=0.0
		4	15/120=0.13	1/16=0.07	0/12=0.0	0/16=0.0	0/17=0.0
		5	19/122=0.16	2/18=0.12	3/19=0.16	2/17=0.12	1/18=0.06
		6	13/136=0.10	0/18=0.0	2/11=0.19	0/18=0.0	1/13=0.08
	•	7	16/135=0.12	2/19=0.11	0/ 9=0.0	1/12=0.09	0/12=0.0

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT RELATIONSHIPS AND DIFFERENCES USING THE HISTORICAL CONTROL GROUP

ONE !, * = SIGNIFICANT AT P LESS THAN 0.05
TWO !.* = SIGNIFICANT AT P LESS THAN 0.01

^{*} SIGNIFICANTLY DIFFERENT FRCM CONTROL

[!] SIGNIFICANT LINEAR RELATIONSHIP WITH ARITH OR LOG DOSE (HEADING OF COLUMN)

TABLE VIII

COMPOUND 36

STUDY SUBACUTE

DEAD IMPLANTS / TOTAL IMPLANTS

WFEK	HISTORICAL CONTROL	NEGATIVE CONTROL	DOSE LEVEL 50.000 MG/KG	DOSE LEVEL 500.000 MG/KG	DOSE LEVEL 5000.000 MG/KG
1	40/1231=0.04	8/167=0.05	5/150=0.04	3/108=0.03	9/163=0.06
2	59/1474=0.05	18/190=0.10	33/178=0.19	49/222=0.23 *ā	•
3	69/1405=0.05	3/139=0.03	4/155=0.03	0/181=0.0	1/181=0.01 **aaD **aaD
4	66/1414=0.05	7/191=0.04	3/137=0.03	3/180=0.02	0/195=0.0 *ap
5	78/1462=0.06	7/219=0.04 aD	7/237=0.03	12/199=0.07	7/231=0.04 *ap
6	62/1626=0.04	1/179=0.01 *aap	15/108=0.14	0/183=0.0	4/132=0.04
7	70/1566=0.05	5/204=0.03	0/100=0.0	4/137=0.03	1/139=0.01 **@@D

SYMBOLS ON FIRST LINE DENOTE SIGNIFICANT DIFFERENCES USING THE NEGATIVE CONTROL GROUP

SYMBOLS ON SECOND LINE DENOTE SIGNIFICANT DIFFERENCES USING THE HISTORICAL CONTROL GROUP

- * = TWO-TAILED TEST
- @ = ONE-TAILED TEST
- ONE *, $\hat{\sigma}$ = SIGNIFICANT AT P LESS THAN 0.05 TWO *, $\hat{\sigma}$ = SIGNIFICANT AT P LESS THAN 0.01
- *, & SIGNIFICANTLY DIFFERENT FROM CONTROL

APPENDICES

II. MATERIALS AND METHODS

A. <u>Animal Husbandry</u>

1. Animals (Rats and Mice)

Ten to twelve week old rats (280 to 350 g) and male mice (25 to 30 g) were fed a commercial 4% fat diet and water ad libitum until they were put on experiment. Flow Laboratories random-bred, closed colony, Sprague-Dawley CD strain rats were used in the cytogenetic studies. Flow Laboratories ICR male mice were employed in the Host-Mediated Assay.

2. Preparation of Diet

A commercial 4% fat diet was fed to all animals. Periodic tests to verify the absence of coliforms, <u>Salmonella</u> and <u>Pseudomonas</u> sp. were performed.

3. Husbandry

Animals were held in quarantine for 4-11 days. Mice were housed five to a cage and rats one to five to a cage. Animals were identified by ear punch. Sanitary cages and bedding were used, and changed two times per week, at which time water containers were cleaned, sanitized and filled. Once a week, cages were repositioned on racks; racks were repositioned within rooms monthly. Personnel handling animals or working within animal facilities wore head coverings and face masks, as well as suitable garments. Individuals with respiratory or other overt infections were excluded from the animal facilities.

B. <u>Dosage Determination</u>

1. Acute LD_{50} and LD_{5} Determination Since the compounds proposed for testing are included in



the food additive regulations as "generally recognized as safe" (GRAS), it was expected that a large number of them would be sufficiently non-toxic so that determination of a LD_{50} or a LD_{5} would be of no practical value. In fact, this has been our experience with previously tested compounds from this list. In the case of these relatively non-toxic compounds, attempts were made to assure that the amounts to be administered would not affect the animals by means (mechanical, physical, etc.) related to their bulk rather than to their toxicity. In the cases of certain compounds where a LD_{50} or a LD_{5} could not be determined, an exceedingly high concentration, 5 g/kg, was employed and accepted as the LD_{5} level. In cases where the toxicity was high enough to allow determination of a LD_{5} , the following protocol was used.

Thirty rats of the strain chosen for studies described below and of approximately the age and weight specified were assigned at random to six groups. Each group was then given, using the chosen route of administration, one of a series of dosages of the test compound following a logarithmic dosage scheme. The series of dosages were derived from a consideration of whatever toxicity information was available for the particular test compound. The objective in selecting dosages was to choose values which would cause mortalities between 10% and 90%.

When information was inadequate to derive a suitable series of dosages, five rats were used to identify the proper range. Each of these was given one of a widely spaced (differing by 10X) series of doses. This was confidently expected to suffice for derivation of the series of dosages to be used in the LD_{50} determination.



The mortalities observed when the series of dosages were given to the 30 rats were then subjected to a probit analysis and calculation of LD_{50} , LD_{5} , slope and confidence limits by the method of Litchfield and Wilcoxon. The highest dose level used was either a finite LD_{5} or 5000 mg/kg. The intermediate level used was either 1/10 of the finite LD_{5} or 2500 mg/kg. The low level used was either 1/100 of the finite LD_{5} or 30 mg/kg.

2. Subacute Studies

Subacute doses were identical to those used in the acute studies. Each subacute study animal was given the acute dosage once a day for each of five consecutive days (24 hours apart).

C. <u>Mutagenicity Testing Protocols</u>

Host-Mediated Assay

Flow Laboratories ICR random-bred male mice were used in this study. In the acute and subacute studies ten animals, 25-30 g each, were employed at each dose level. Solvent and positive controls were run at all times. The positive control (dimethyl nitrosamine) was run by the acute system only at a dose of 100 mg/kg for Salmonella. For yeast, ethyl methane sulfonate (EMS) intramuscularly injected at a dose of 350 mg/kg was used. The solvents used and the toxicity data are presented in the Results and Discussion Section of the report.

The indicator organisms used in this study were: (1) two histidine auxotrophs (his G-46, TA-1530) of <u>Salmonella typhimurium</u>, and (2) a diploid strain (D-3) of <u>Saccharomyces cerevisiae</u>. The induction of reverse mutation was determined with the <u>Salmonella</u>; mitotic recombination was determined with yeast. Chemicals were evaluated directly by <u>in vitro</u> bacterial and yeast studies prior to, or concurrent with, the studies in



mice. Only animals on the subacute studies were not fed the evening prior to compound administration. The Salmonella were carried in tryptone yeast extract gel, transferred weekly. They were transferred to tryptone yeast extract broth 48 hours before use: they were transferred a second time from broth to broth 24 hours prior to use, and again 8 hours before use. The mouse inoculum was prepared by transferring 4 ml of the 8-hour broth culture to 50 ml broth bottles which had been prewarmed at 37°C. Exponential log-phase organisms were inoculated intraperitoneally into the mice approximately 2-1/2 hours later when the appropriate density indicating 3.0 \times 10^8 cells/ml was reached. The Saccharomyces was carried in yeast complete agar. The inoculum was prepared by harvesting the organisms from the surface of the plates with sterile saline. The cells were washed three times with sterile saline and suspended in a concentration of 5.0 \times 10⁸ cells/ml. Two ml of the suspension was inoculated into each mouse intraperitoneally. Total plate counts on Salmonella were on tryptone yeast extract and for Saccharomyces on yeast complete medium.

a. Acute study

Three dosage levels (usage, intermediate [determined as discussed previously], and LD $_5$) were administered orally by intubation to ten mice. Positive controls and negative vehicle controls were included in each study. All animals received 2 ml of the indicator organism intraperitoneally. Each ml contained 3.0 x 10^8 cells for Salmonella and 5.0 x 10^8 cells for Saccharomyces. Three hours later, each animal was killed and 2 ml of sterile saline was introduced intraperitoneally. As much fluid as possible was then aseptically removed from the peritoneal cavity. Dilution blanks for bacteria containing 4.5 ml of serile saline were prepared in advance. Tenfold serial

dilutions were made of each peritoneal exudate (0.5 ml exudate + 4.5 ml saline) yielding a concentration series from 10^0 (undiluted peritoneal exudate) through 10^{-7} . For enumeration of total bacterial counts, the 10^{-6} and 10^{-7} dilutions were plated on tryptone yeast extract agar, 3 plates/sample, 0.2 ml sample/ plate. Each sample was spread over the surface of the plate using a bent glass rod immersed in 95% ethanol and flamed just prior to use. In plating for the total mutant counts on minimal agar, the 10^0 dilution was used, 0.2 ml being plated on each of 5 plates. The plating procedure was identical to that followed for the tryptone yeast extract agar plates. All plates were incubated at 37°C, tryptone yeast extract agar plates for 18 hours and minimal agar plates for 40 hours. For yeast mitotic recombination, dilution blanks containing 4.5 ml of sterile saline were prepared in advance. Tenfold serial dilutions were made of each sample yielding a series from 10^{0} to 10^{-5} . Samples of 0.1 ml of the 10^{-5} , 10^{-4} , and 10^{-3} dilutions were removed and plated on complete medium (10 plates each). All plates were incubated at 30° C for 40 hours. The 10^{-5} dilutions were used to determine total populations and the 10^{-4} and 10^{-3} plates were examined after an additional 40 hours at 4°C for red sectors indicating a mutation. Bacterial scoring was calculated as follows:

Total mutants on 5 plates x appropriate exponent = CFU/ml (CFU is Colony Forming Units) of sample plated CFU/ml x one/dilution factor ($10^{0} - 10^{-7}$) = CFU/ml in undiluted exudate. The mutation frequency (MF) calculated for each sample was:

 $MF = \frac{\text{total mutant cells}}{\text{total population}}$

 $MFt/MFc = \frac{MF \text{ of experimental sample}}{MF \text{ of control sample}}$

(MFt/MFc = 1.00 for control sample)



Yeast mitotic recombinants (presumptive <u>ade 2</u>, <u>his 8</u> homozygotes) were seen as red colonies or as red sectors on a normally white yeast colony. The plates (from 10^{-4} and 10^{-3} dilutions) were scanned under the 10X lens of a dissecting scope to enumerate the red colonies and sectors. Population determinations were made from the 10^{-5} dilution plates. A recombinant frequency (RF) was calculated:

RF = total recombinants counted total number colonies screened

b. Subacute study

Similar groups of animals at each dose level received five oral doses of the test compound 24 hours apart. Within 30 minutes after the last dosing, the animals were inoculated with the test organism and handled in the same fashion as those in the acute study.

c. <u>In vitro</u> study

Cultures of <u>S. typhimurium</u> histidine auxotrophs

(G-46 and TA-1530) were plated on appropriate media. The test compound was then added to the plate, either in the form of a microdrop of solution (0.01 to 0.25 ml) applied to a small filter paper disc resting on the agar or a small crystal applied directly to the agar. Tenfold serial dilutions of the culture were employed and plated so as not to miss the optimum cell density for mutant growth. Mutant colonies were observed and scored. Strain D-3 <u>Saccharomyces</u> cells at proper dilutions were shaken with the test compound, diluted, and plated at 50% survival level or above (see HMA Supplementary Materials and Methods). Red sectors were then scored and the frequency calculated after suitable incubation. Negative and positive controls were run concurrently. The positive control was EMS for <u>Salmonella</u> and <u>Saccharomyces</u>. The <u>in vitro Salmonella</u> tests were reported



as (+) or (-) or questionable; the <u>in vitro Saccharomyces</u> tests were reported as sample concentrations, percent survival, and recombinants/ 10^5 survivors. For the <u>Saccharomyces</u> a 50% survival level, e.g., an arbitrary 5.0% w/v test level, was used when no LD₅₀ was determinable.

2. Cytogenetic Studies

a. <u>In vivo</u> study

Ten to twelve week old, male, albino rats obtained from a closed colony (random-bred) were used. A total of 59 animals in the acute study and 18 animals in the subacute study was used, as illustrated in the following protocol.

Number of Animals Used

Acute Study

<u>Treatment</u>	Time Killed	d After Admir	nistration
	6 Hours	24 Hours	48 Hours
High Level	5	5	. 5
Intermediate Level	5	5	5
Low Level	5	5	5
Positive Control	0	0	5
Negative Control	3	. 3	3

Subacute Study

Five doses 24 hours apart; animals killed 6 hours after last dose.

Treatment	Killed After Administration
High Level	5
Intermediate Level	5
Low Level	5
Negative Control	3

All animals were dosed by gastric intubation.

Four hours after the last compound administration, and two hours prior to killing, each animal was given 4 mg/kg of colcemid intra-



peritoneally in order to arrest the bone marrow cells in C-mitosis. Animals were killed by using CO₂, and the adhering muscle and epiphysis of one femur were removed. The marrow "plug" was removed with a tuberculin syringe and an 18 gauge needle, aspirated into 5 ml of Hanks' balanced salt solution (BSS) in a test tupe and capped. The specimens were centrifuged at 1,500 RPM in a table-top centrifuge for 5 minutes, decanted, and 2 ml of hypotonic 0.5% KCl solution was added with gentle agitation to resuspended the cells. The specimens were then placed in a 37°C water bath for 20 minutes in order to swell the cells. Following centrifugation for 5 minutes at 1,500 RPM, the supernatant was decanted and 2 ml of fixative (3:1 absolute methanol:glacial acetic acid) was added. The cells were resuspended in the fixative with gentle agitation, capped, and placed at 4°C for 30 minutes. The specimens were again centrifuged, decanted, 2 ml of prepared fixative was added, and the cells were resuspended and placed at 4°C overnight.

The following day the specimens were again centrifuged, decanted and 0.3 - 0.6 ml of freshly prepared fixative was added to obtain a suitable density. The cells were resuspended and 2 - 3 drops of the suspension were allowed to drop onto a clean, dry slide held at 15° from the horizontal. As the suspension flowed to the edge of the slide, it was ignited by an alcohol burner and allowed to flame. Following ignition, the slides were allowed to dry at room temperature overnight. Duplicate slides were prepared. The slides were stained using a 5% Giemsa solution (Giemsa buffer pH 7.2) for 20 minutes, rinsed in acetone, 1:1 acetone:xylene, and placed in fresh xylene for 30 minutes. The slides were then mounted using Permount (Fisher Scientific) and 24 x 50 mm coverglasses. The coverglasses were selected to be 0.17 mm \pm 0.005 mm in thickness by use of a coverglass micrometer. The preparations



were examined using Leitz Ortholux I & II microscopes with brightfield optics and xenon light sources. These specimens were scanned with 10X and 24X objectives and suitable metaphase spreads that were countable were then examined critically using 40X, 63X or 100X oil immersion flatfield apochromatic objectives. Oculars were either 12X or 16X widefield periplanatics and the tube magnification either 1X or 1.25X. The filters used were either a didymium (BG20) or a Schott IL570 m μ interference filter.

The chromosomes of each cell were counted and only diploid cells were analyzed. They were scored for chromatid gaps and breaks, chromosome gaps and breaks, reunions, cells with greater than ten aberrations, polyploidy, pulverization, and any other chromosomal aberrations which were observed. They were recorded on the currently used forms and expressed as percentages on the summary sheets. Fifty metaphase spreads were scored per animal. Mitotic indices were obtained by counting at least 500 cells and the ratio of the number of cells in mitosis/the number of cells observed was expressed as the mitotic index.

Positive controls in the acute study consisted of animals which had been given the known mutagen Triethylene Melamine (TEM) administered intraperitoneally at a level of 0.30 mg/kg. Negative controls on the acute and subacute studies consisted of the vehicle in which the compound was administered. The dosage levels, solvents and toxicity data are included in the Results and Discussion Section of the report.

b. <u>In vitro</u> study

Human embryonic lung cultures (WI-38) which were negative for adventitious agents (viruses, mycoplasma) which may interfere



were used. These cells were employed at passage level 19. The cells had been transferred using 0.025% trypsin and planted in 32 oz. prescription bottles containing 40 ml of tissue culture medium. When growth was approximately 95% confluent the cells were removed from the glass using trypsin, centrifuged, and frozen in tissue culture medium containing dimethyl sulfoxide (DMSO). Cells were frozen in vials in the vapor phase of liquid nitrogen at a concentration of 2 \times 10^6 cells/ml. When needed, the vials were removed from liquid nitrogen, quick-thawed in a 37°C water bath, washed free of DMSO, suspended in tissue culture medium (minimal essential medium [MEM] plus 1% glutamine, 200 units/ml of penicillin and 200 µg/ml of streptomycin and 15% fetal calf serum) and planted in milk dilution bottles at a concentration of 5 x 10^5 cells/ml. The test compound was added at three dose levels using three bottles for each level, 24 hours after planting. The dose levels required a preliminary determination of a tissue culture toxicity. This was accomplished by adding logarithmic doses of the compound in saline to a series of tubes containing 5 x 10^5 cells/ml which were almost confluent. The cells were examined at 24, 48, and 72 hours. Any cytopathic effect (CPE) or inhibition of mitoses was scored as toxicity. Five more closely spaced dose levels were employed within the two logarithmic dosages, the higher of which showed toxicity and the lower no effect. The solvents used and the range finding data are presented in the toxicity data report under Results and Discussion. The dose level below the lowest toxic level was employed as the high level. Logarithmic dose levels were employed for the medium and low levels.

Cells were incubated at 37°C and examined twice daily to determine when an adequate number of mitoses were present. Cells were harvested by shaking when sufficient mitoses were observed, usually 24 - 48



hours after planting, centrifuged, and fixed in absolute methanol:glacial acetic acid (3:1) for 30 minutes.

The specimens were centrifuged, decanted, and suspended in acetic acid-orcein stain (2.0%) and a drop of suspension placed on a clean dry slide. Selected coverglasses 0.17 mm in thickness were placed on the suspension and the excess stain gently expressed from the slide. The coverglasses were sealed with clear nail polish and examined immediately.

The microscopes, objectives, oculars, filters and light sources were enumerated under the metaphase description. Positive controls used were TEM (at a concentration of 0.1 mcg/ml dissolved in saline) and negative controls which consisted of the vehicle in which the test compound was dissolved, which was 0.85% saline. Data were reported on forms currently used and expressed as percentages on the anaphase summary sheets.

3. Dominant Lethal Assay

In this test, male and female random bred rats from a closed colony were employed. These animals were 10-12 weeks old at the time of use. Ten male rats were assigned to each of 5 groups; 3 dose levels selected as described above, a positive control (triethylene melamine) (TEM) and a negative control (solvent only). The positive control was administered intraperitoneally. Administration of the test compound was orally by intubation in both the acute study (1 dose) and in the subacute study (1 dose per day for 5 days). Following treatment, the males were sequentially mated to 2 females per week for 8 weeks (7 weeks in the subacute study). Two virgin female rats were housed with a male for 5 days (Monday through Friday). These two females were removed and housed in a cage until killed. The male was rested on Saturday and Sunday and two new females introduced to the cage on



Monday. It has been our experience that conception has taken place in more than 90% of the females by Friday and that the two day rest is beneficial to the male as regards subsequent weekly matings. Females were killed using ${\rm CO}_2$ at 14 days after separating from the male, and at necropsy the uterus was examined for deciduomata (early deaths), late fetal deaths and total implantations.

Sufficient animals were provided in our experimental design to accommodate for any reduction in the number of conceptions. Each male was mated with two females per week, and this provided for an adequate number of implantations per group per week (200 minimum) for negative controls, even if there was a fourfold reduction in fertility of implantations. Results were analyzed according to the statistical procedures described in Supplementary Materials and Methods. Corpora lutea, early fetal deaths, late fetal deaths and total implantations per uterine horn were recorded on the raw data sheets, which are submitted separately.

- D. Supplementary Materials and Methods
 - 1. Host-Mediated Assay <u>In Vitro</u> and Formulae
 - a. Bacterial in vitro plate tests

This method has been published by Ames: The Detection of Chemical Mutagens with Enteric Bacteria, in <u>Chemical Mutagens</u>; <u>Principles and Methods for Their Detection</u>, Vol. 1, Chapter 9, pp. 267-282, A. Hollaender, Editor, Plenum Press, New York (1971).

- b. <u>In vitro</u> for mitotic recombination
- (1) Strain D-3 was grown to stationary phase on complete medium agar plates at 30°C (3-4 days). Cells were rinsed from the plates and washed twice in saline and cell concentration determined spectro-



photometrically. (A standard curve previously determined for colony forming units versus % transmittance at 545 mu was easily used.)

- (2) Cells from the concentration suspension were diluted appropriately into 0.067 M Phosphate buffer pH 7.2 to provide 5×10^7 cells/ml in a total of 25 ml.
- 4 hours at 30°C, with shaking, at concentrations which permitted determination of the 50% survival level. Then, if not included in the first experiment, the compound was tested again only at the 50% survival level. If 50% survival level could not be determined, the arbitrary test level of 5% w/v was used.
- plated on complete agar medium for determination of total population and red sectors. Total surviving population was conveniently measured on plates of 10^{-4} and 10^{-5} dilutions using 0.2 ml per plate (5 plates), and sectors determined on plates of 10^{-3} and 10^{-4} dilutions using 0.2 ml per plate (5 plates). Plates were incubated for 2 days at 30°C followed by a holding period of 2 days at 4°C to promote color development with limited enlargement of the colonies. Red sectors were scored by systematically scanning the plates with a dissecting microscope at 10X magnification.
- (5) The frequency of red sectors can then be calculated and may be expressed conveniently as sectors per 10^5 survivors for comparison with untreated controls.
- (6) Ethyl Methane Sulfonate (EMS) was employed as the positive control in both <u>in vitro</u> systems.
 - c. Minimal medium (bacteria):
 Spizizen's Minimal Medium:



4X Salt Solution:

(NH₄) SO₄

8.0 gm

 K_2HP0_4

56.0 gm

KH2PO4

24.0 gm

Na Citrate

4.0 gm

Mg SO₄

0.8 gm

Biotin

0.004 gm

H₂0

qs to 1 liter

Sterilize by autoclaving

(121°C/15 min.)

<u>Medium</u>:

4X Salt Solution

:250 ml

5.0% Glucose (sterile)

:100 ml (If histidine is added

at concentration of 30 mg/liter, this becomes a complete bacterial

medium.)

1.5% Bacto-agar (sterile)

:650 ml

d. Complete medium (bacteria):

Bacto-Tryptone

1.0 gm

Yeast-Extract

0.5 gm

Bacto-Agar

2.0 gm

Distilled H₂0

100.0 ml

Sterilize by autoclaving (121°C for 15 minutes).

e. Complete medium (yeast):

KH2P04

1.5 gm

 $MgSO_4$

0.5 gm

 $(NH_4)_2SO_4$

4.5 gm

 Peptone
 3.5 gm

 Yeast-Extract
 5.0 gm

 Glucose
 20.0 gm

 Agar
 20.0 gm

 Distilled H20
 1000.0 ml

Sterilize by autoclaving (121°C for 15 minutes).

 Cytogenetics <u>In Vitro</u> Preparation of Anaphase Chromosomes (from Nichols, 1970)

"Anaphase preparations may be made by several methods. convenient approach is to grow cells directly on coverslips in petri dishes. With human fibroblasts 400,000 cells added to a 22 \times 44 mm coverslip in a 50 mm petri dish grown in a 5% ${\rm CO}_2$ atmosphere in air has proved very satisfactory. When adequate numbers of mitoses are visualized directly utilizing an inverted microscope (usually 48 to 92 hours after planting) the coverslip is transferred to absolute ethanol for 15 minutes for fixation. They are then stained with any one of a number of suitable stains (Fuelgen, May-Grunwald-Giemse, orcein) and attached to a slide with mounting media for evaluation. Anaphase preparations may also be prepared on cells grown in suspension or cells from a monolayer that have been put into suspension. In this instance the cells are centrifuged and fixed with the squash fixative. They are then suspended in the stain and a drop of the suspension put on the slide and covered with a coverslip. However, in this case, only the excess stain is gently expressed from under the coverslip and no squashing is carried out. In anaphase preparations no pretreatment with colchicine or hypotonic expansion is used and no technique for spreading the cells is used, so that the spindle and normal relationships of the chromosomes are not disturbed."



- 3. Statistical Analyses of Dominant Lethal Studies

 The following statistical analyses were employed as a means of analyzing the results of the dominant lethal studies.
 - a. The fertility index

The number of pregnant females/number of mated females with the chi-square was used to compare each treatment to the control. Armitage's trend was used for linear proportions to test whether the fertility index was linearly related to arithmetic or log dose.

b. Total number of implantations

The t-test was used to determine significant differences between average number of implantations per pregnant female for each treatment compared to the control. Regression techniques were used to determine whether the average number of implantations per female was related to the arithmetic or log dose.

- c. Total number of <u>corpora lutea</u>

 The t-test was used to determine significant differences between average number of <u>corpora lutea</u> per pregnant female for each treatment compared to the control.
 - d. Preimplantation losses

Preimplantation losses were computed for each female by subtracting the number of implantations from the number of corpora lutea. Freeman-Tukey transformation was used on the preimplantation losses for each female and then the t-test was used to compare each treatment to control. Regression technique was used to determine whether the average number of preimplantation losses per female was related to the arithmetic or log dose.



e. Dead implants

Dead implants were treated the same as pre-

implantation losses.

f. One or more dead implants

The proportion of females with one or more dead implants was computed, each treatment compared to control by chi-square test and Armitage's trend used for linear proportions to see if proportions were linearly related to either arithmetic or log dose. Also, probit regression analysis was used to determine whether the probit of the proportions was related to log dose.

g. Two or more dead implants

The proportion of females with two or more dead implants computed was treated same as above (f).

h. Dead implants per total implants

Dead implants per total implants were computed for each female and used Freeman-Tukey arc-sine transformation on data for each female; then used t-test to compare each treatment to control.

Historical control data was compiled on a continuous basis as studies were completed. In addition to comparing each treatment to control, as outlined above, each treatment was compared to a historical control.

In order to take variation between males into account, a nested model was used. An analysis of across weeks is also provided.

In addition to these tests, the distribution forms of the various parameters were tested in order to evaluate the appropriateness of some of the tests being used. Certain correlations between parameters may exist and were examined as one step to determine the appropriateness of models. If necessary, alternate test methods were implemented.



The results are presented in tabular form with the addition of historical control information. In addition to these tables, a written report of all findings is provided. As information became available from the on-going investigation of these data, it was reported and suggestions included for changes to the methods of analysis. The statistical reports give the level of significance using both a one-tailed and two-tailed test. Finally, a summary sheet for each study is provided.



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Males are randomly drawn from infinite population

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F. Abbreviations

- 1. mu = micron
- 2. mcg = ug = microgram
- 3. g = gram
- 4. kg = kilogram
- 5. ml = milliliter
- 6. rpm = revolutions per minute
- 7. °C = degrees centigrade
- 8. pH = power of the hydrogen ion concentration to the base 10
- 9. M = molar solution
- 10. conc. = concentration
- 11. MTD = maximum tolerated dosage = High = LD_5 if determined or else exceedingly high dose, such as 5 g/kg
- 12. INT = intermediate = medium level
- 13. USE = usage level if known = low level
- 14. BSS = balanced salt solution
- 15. C-metaphase = cells arrested in metaphase, using colchine or colcemid
- 16. LD_{50} = that dosage which produced 50% mortality in the group of animals treated
- 17. LD₅ = that dosage which produced 5% mortality in the group of animals treated
- 18. NC = negative control
- 19. PC = positive control
- 20. AU = acute usage level (low level)
- 21. AI = acute intermediate level (medium level)
- 22. AMTD = acute maximum tolerated dose level (LD_5 level, high level)



- 23. SAU = subacute usage level (low level)
- 24. SAI = subacute intermediate level (medium level)
- 25. SA LD_5 = subacute LD_5 level (MTD level, high level)
- 26. CO_2 = carbon dioxide
- 27. DMN = Dimethyl nitrosamine
- 28. EMS = Ethyl methane sulfonate
- 29. TEM = Triethylene melamine
- 30. DMSO = Dimethyl sulfoxide
- 31. MEM = minimal essential medium (Eagle's)
- 32. CPE = cytopathic effect
- 33. his = histidine marker
- 34. D-3 = mitotic recombinant strain of Saccharomyces
- 35. mf = mean mutant frequency
- 36. MFt/MFc = mean mutant frequency of the test compound group compared to mean mutant frequency of the negative control group
- 37. CFU = colony forming units
- 38. WI-38 = code name for a strain of human embryonic lung tissue culture cells
- 39. Rec x 10^5 = mitotic recombinants x 10^5
- 40. Mean B/A = mean frequency
- 41. tot. scr. = total scored
- 42. tot. = total
- 43. χ^2 = a test of variation in the data from the computed regression line tested in these studies at the 5% level
- 44. Aber. = aberrations
- 45. Frag. = fragment
- 46. HMA = host-mediated assay

